

Educational Proceedings Book

August 11-12, 2017 Santiago, Chile



INTERNATIONAL MYELOMA SOCIETY

Educational Workshops

Santiago, Chile-August 11-12, 2017 (Crowne Plaza)
Prague, Czech Republic-September 29-30, 2017 (Marriott)
National Harbor, Maryland-October 27-28, 2017 (Gaylord National Hotel)

The International Myeloma Society (IMS) is a professional, scientific and medical society established to bring together clinical and experimental scientists involved in the study of myeloma. The IMS accomplishes its mission by providing educational forums in the form of conferences, symposia, and workshops conducted around the globe.

The goal for the Educational Symposia Series is to share with local hematologists the **state of the art on diagnosis, monitoring and treatment** of Plasma Cell malignancies paying special attention to the *current practice in the local countries* and the opportunities for improvement. Our purpose is also to offer the opportunity for **social interaction** in order to establish potential *collaborations* with the invited speakers including the possibility of visiting their institutions or establishing fellowship programs.

The program will include **lectures plus roundtable discussions around the most hot and controversial topics**. The format of the roundtable will include the *counterpoint opinion* of the invited speakers with the local hematologists based on both challenging questions and clinical cases.

Local program chair(s) are selected to help to ensure the content will address the needs from the regions/countries in which the programs are held.

Santiago Program Chairs:

Guilhermo Conte, MD Hospital Clinico de la Universidad de Chile

Vania Hungria, MD Irmandade da Santa Clara de Misericordia de Sao Paulo

Dorotea Fantl, MD Hospital Italiano de Buenos Aires

Prague Program Chairs:

Roman Hajek, MD University Hospital Ostrava

Ivan Spicka, MD VHN Charles University Prague

Jiri Minarik, MD University Hospital, Olomouc

SANTIAGO, CHILE

PRELIMINARY PROGRAM

This activity has been planned and implemented in accordance with the accreditation requirements and policies of the Accreditation Council for Continuing Medical Education (ACCME) through the joint providership of Mayo Clinic College of Medicine and Science and the International Myeloma Society. Mayo Clinic College of Medicine and Science is accredited by the ACCME to provide continuing medical education for physicians.

The Mayo Clinic College of Medicine and Science designates this live activity for a maximum of 10.00 AMA PRA Category 1 Credit(s)™. Physicians should claim only the credit commensurate with the extent of their participation in the activity.

Friday, August 11, 2017

07:45-8:00	Welcome and introduction
08:00-08:30	The myeloma pathogenesis with focus on the immune system Ivan Borrelo, MD
08:30-9:00	Genetics in MM patients Rafael Fonseca, MD
09:00-09:30	What are the optimal Imaging Techniques in MM? MRI, CT-scan and PET? Marivi Mateos, MD
09:30-10:00	Coffee
10:00-10:30	How to perform an appropriate protein screening and decisions on follow-up? <i>Joan Blade</i> , MD
10:30-11:00	Minimal residual Disease. How and when to do it? Jesús San Miguel, MD
11:00-12:00	Roundtable: How to do a correct diagnosis and follow-up, including the new diagnostic and response criteria Moderator: Jesús San Miguel, MD: 2-3 clinical cases Discussants: Guillermo Conte, MD, Dorotea Fantl, MD, Juan Navarro, MD, Joan Bladé, MD, Rafael Fonseca, MD, Marvi Mateos, MD
12:00-13:00	Lunch
13:00-13:30	MGUS & Smouldering Myeloma. How to predict outcome & "To treat or not to treat" Mariví Mateos, MD

Friday, August 11, 2017 (continued)

13:30-15:00 How I treat newly diagnosed transplant candidate MM patients

Sergio Giralt, MD (30').

Roundtable on 6 controversial questions with emphasis in local practice (60') Moderator:

Rafael Fonseca, MD

Discussants: David Gómez-Almaguer, MD, Angelo Maiolino, MD, Natalia Schutz, MD, Joan Blade, MD, Sergio Giralt, MD, Jesus San Miquel, MD

- 1. Best induction
- 2. Early vs. late transplant
- 3. One or two
- 4. Consolidation
- 5. Maintenance
- 6. Allotransplant

15:00-15:30 Coffee

15:30-17:00 How I treat newly diagnosed elderly patients?

Ruben Niesvizky, MD (30')

Roundtable on 5 controversial questions with emphasis in local practice (60')

Moderator: Mariví Mateos, MD

Discussants: Vania Hungria, MD, Jose Luis Lopez, MD, Eloisa Riva, MD, Sergio Giralt, MD, Rafael

Fonseca, MD, Ruben Niesvizky, MD

- 1. Optimal combinations
- 2. Fixed versus continuous therapy
- 3. Do we need alkylators?
- 4. High risk
- 5. Frailty scales

Saturday, August 12, 2017

8:00-9:30 How to make the right choices in the relapsed patient

Enrique Ocio, MD

Roundtable on 5 clinical cases with emphasis in local practice (60')

Moderator: Jesus San Miguel, MD

Discussants: Timoleon Anguita, MD, Edvan Crusoe, MD, Enrique Ocio, MD, Ruben Niesvizky,

MD, Rafael Fonseca, MD

- 1. Biological relapse versus agresive relapse
- 2. Early and late relapse and ASCT
- 3. Triplet versus doublets: is it cost-efective?
- 4. Optimal sequence for new agents
- 5. Allotransplant

9:30-10:00 Amyloidosis: "Under-diagnosed disorder"

Joan Bladé, MD

Saturday, August 12, 2017 (continued)

10:00-10:30 Waldenström Macroglobulinemia

Enrique Ocio, MD

10:30-11:00 Coffee

11:00-11:30 Treatment of Disease Complications

Jesus Berdeja, MD

11:30-12:15 Keynote lecture: Present and future of MM

Nikhil Munshi, MD

12:15 Lunch

THANK YOU TO OUR SPONSORS

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This activity is supported by an educational grant from Celgene Corporation.

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International Myeloma Society Educational Workshop August 2017



6. Once you click **START**, the screen will look similar to the below.

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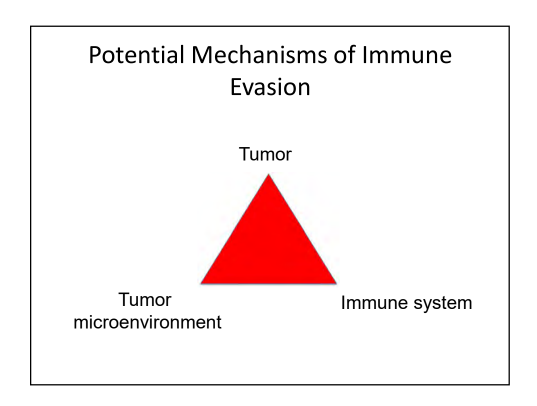
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- Your certificate will remain in your account on <u>ce.mayo.edu</u> under MY ACCOUNT, MY COURSES, TRANSCRIPT.

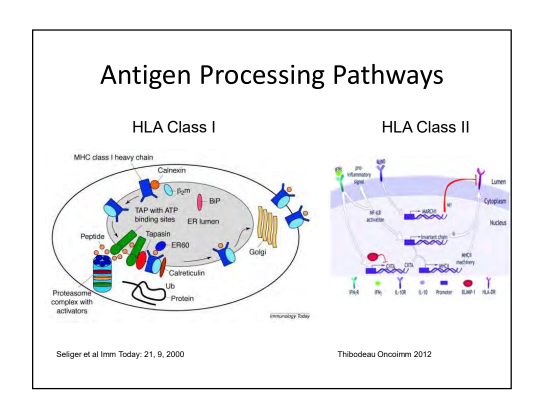
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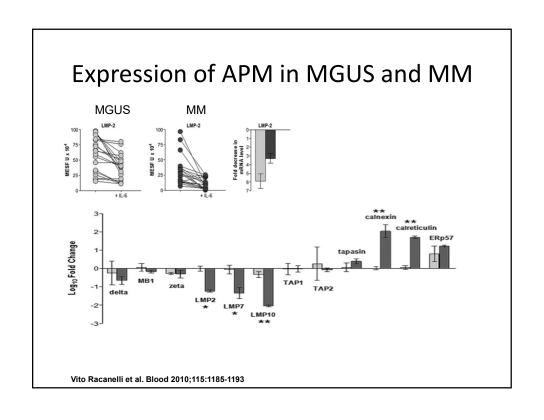
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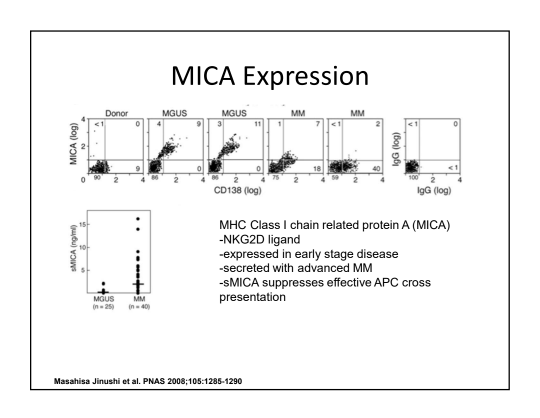
Myeloma Pathogenesis with a Focus on the Immune System

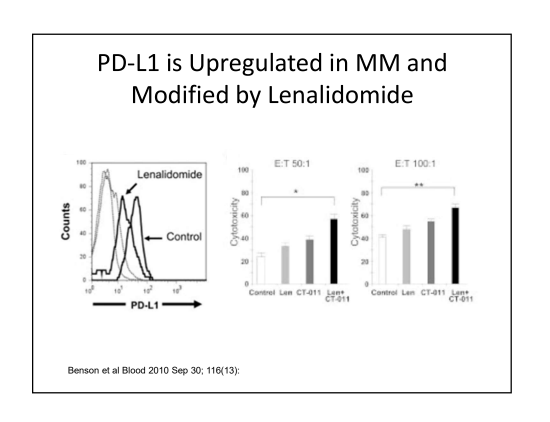
Ivan Borrello, MD Johns Hopkins University

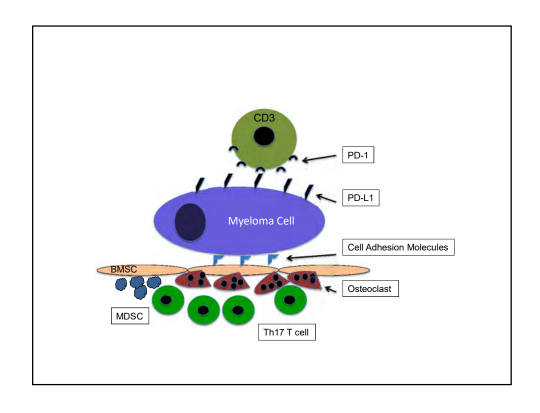


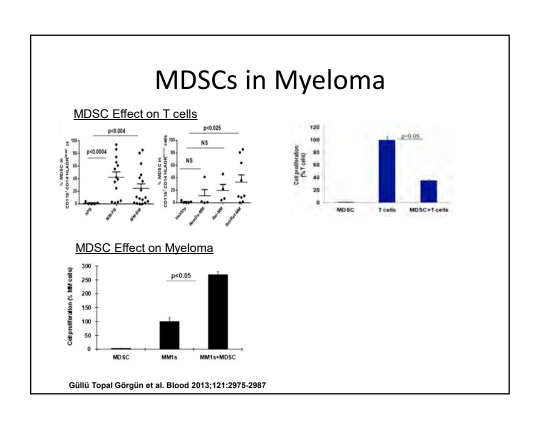


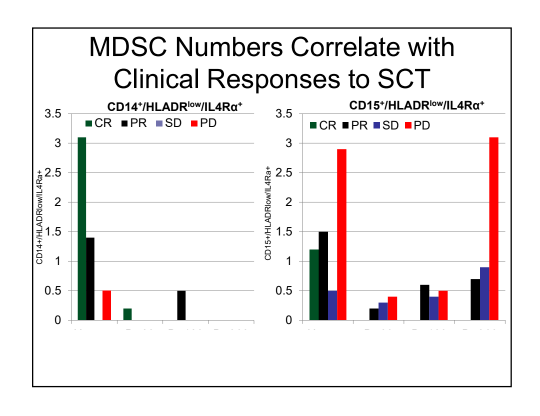


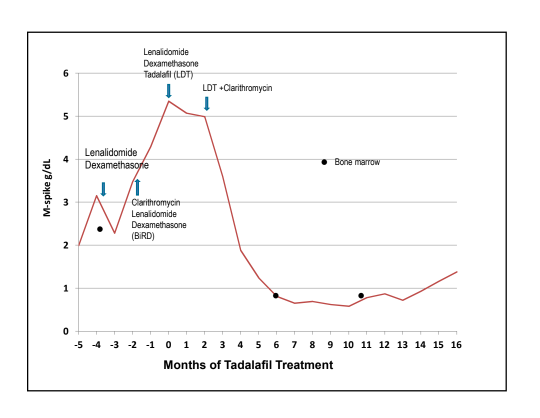




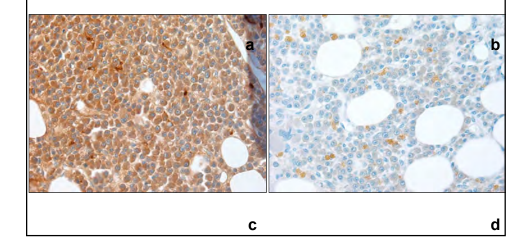




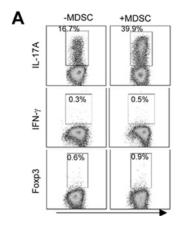




Effect of PDE5 Inhibition on Nitrosylation



MDSCs Induce a Th17 Phenotype

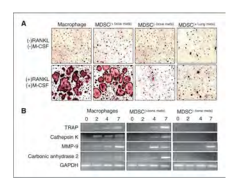


- Autoimmune disease (EAE) is associated with significant expansion of MDSCs
- MDSCs facilitate expansion of Th17 cells

YI, H JI Nov 2012

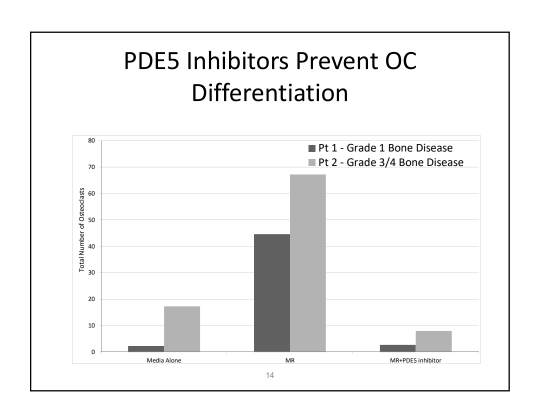
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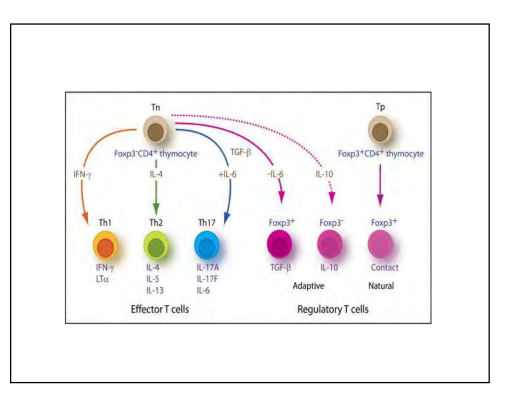
MDSCs from bone metastases induce OC differentiation



- MDSCs from active bone disease show more OC precursors
- Overexpress: iNOS, ROS, HIF1α
- Nitric oxide essential for OC differentiation

Sawant, A Cancer Res. 2013 Jan 15; 73(2): 672-682.



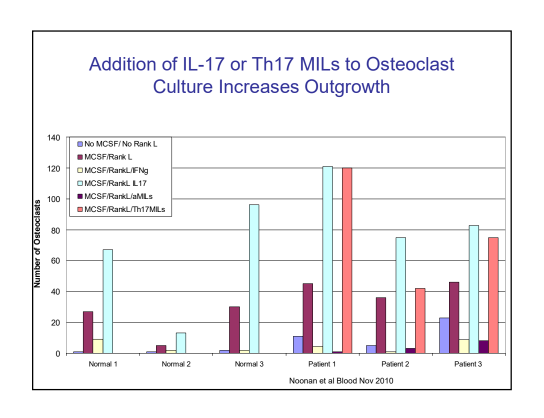


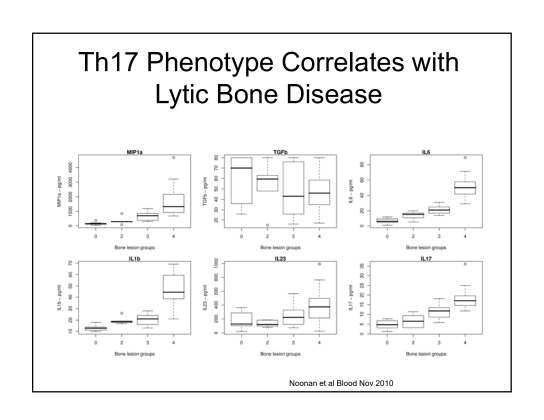
Th17 Profile of Myeloma BM Plasma

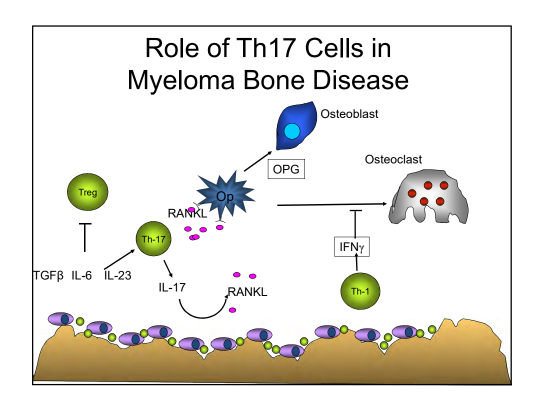
	Myeloma BM	Normal BM	Myeloma PBL	Normal PBL
TGF-β (μg/ml)	59.9	52.6	17.5	16.1
IL-6 (pg/ml)	29.6	3.7	6.5	2.3
IL-23 (pg/ml)	246.4	35.9	19.6	18
IL-17 (pg/ml)	12.2	5.1	0.4	0

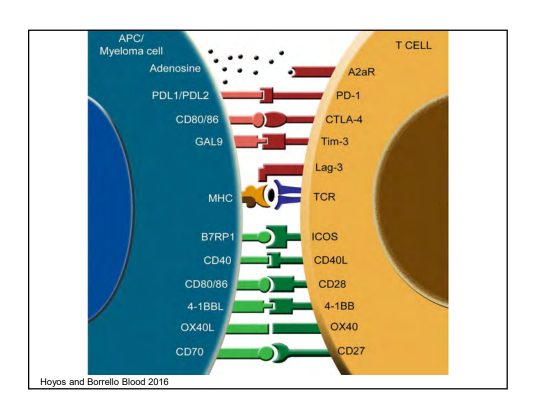
Patient N= 56 Normal N=3

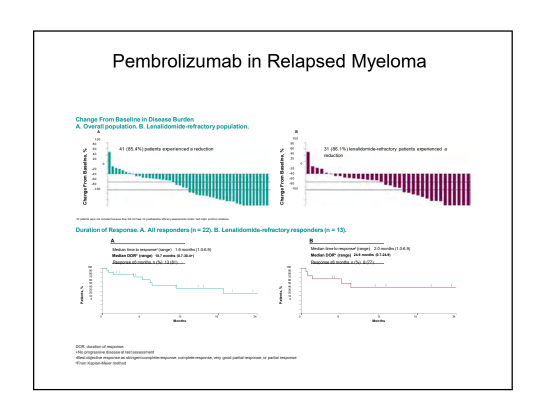
Noonan et al Blood Nov 2010

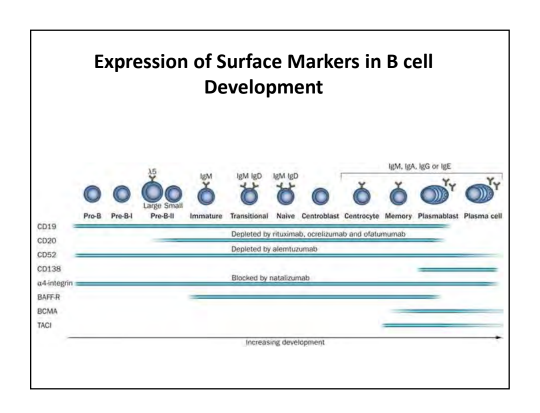


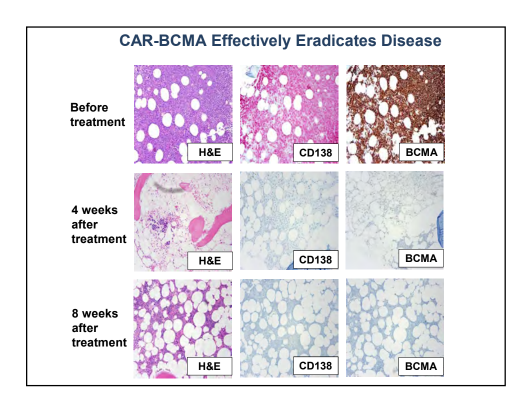












Conclusions

- Immune suppression in tumor microenvironment in myeloma increases from MGUS to active MM
- Mechanisms of immune escape include
 - Downregulation of HLA expression
 - Dysfunction of antigen presentation
 - Increase in MDSCs
 - Development of Th17 cells
 - Increase in PD-L1 expression
 - Upregulation of PD-1 expression on T cells
- Immune therapy can show clinical benefits several of these pathways including:
 - PD-1 blockade
 - CAR-T cells targeting BCMA
 - Vaccines
 - Immunosuppressive pathways: Th17, MDSCs

Rafael Fonseca MD Chair, Department of Medicine Mayo Clinic in AZ

Genetics in Myeloma; The Basics





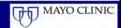


Scottsdale, Arizona

Rochester, Minnesota

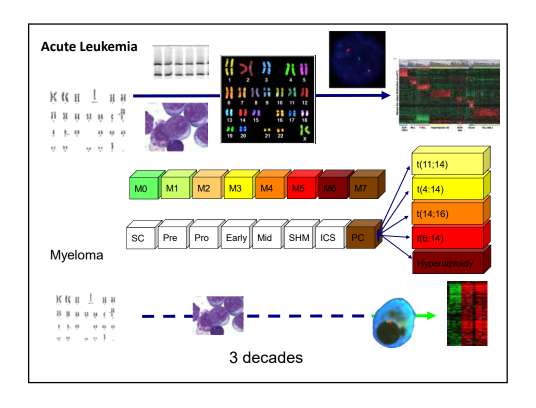
Jacksonville, Florida

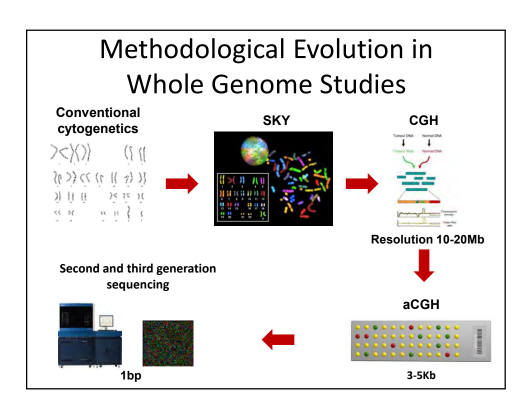
Mayo Clinic College of Medicine Mayo Clinic Comprehensive Cancer Center

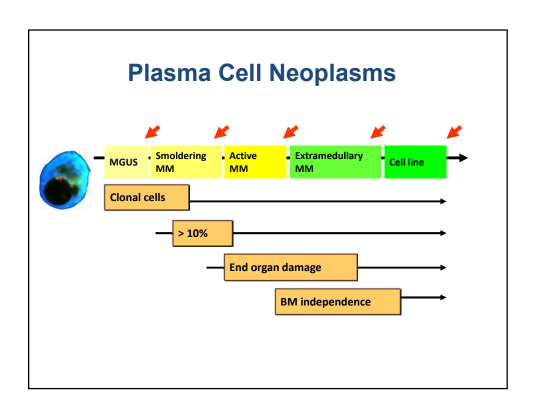


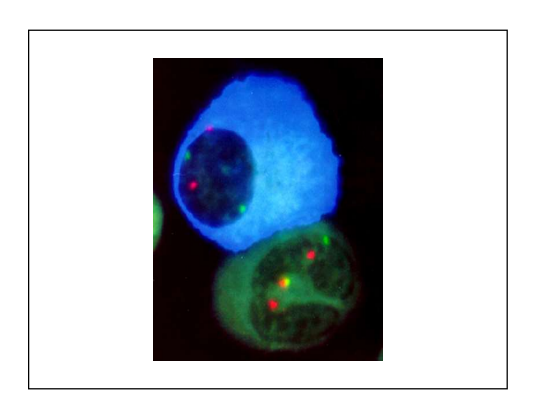
Disclosures

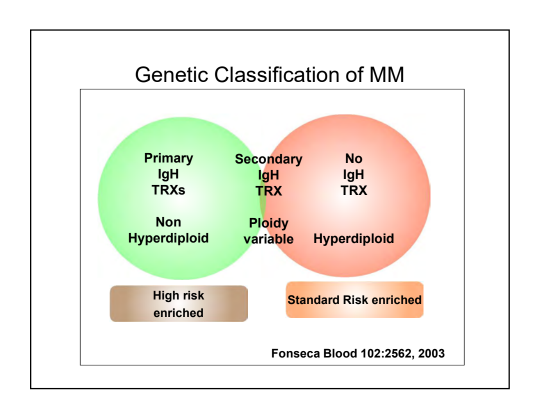
- Consulting: AMGEN, BMS, Celgene, Takeda, Bayer, Jansen, Novartis, Pharmacyclics, Sanofi
- · Speakers Bureaus: None
- SAB: Adaptive Biotechnologies
- Patent for FISH in MM ~\$2000/year
- Registered independent Libertarian
- · Believe in stem cell transplant
- · Dislike wasting your time with this slide

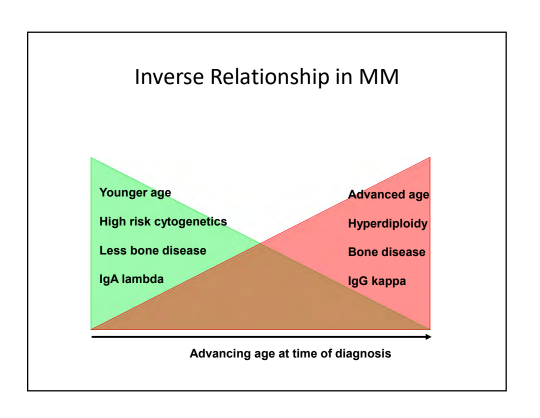


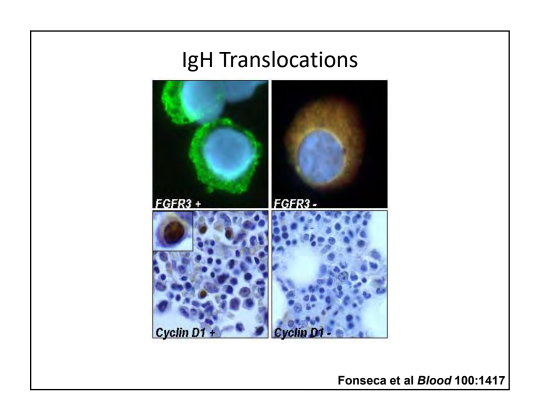


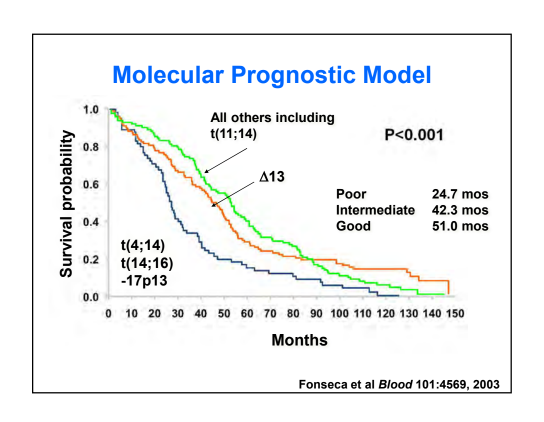


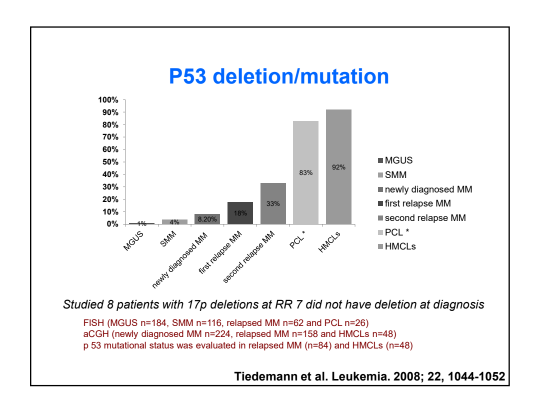






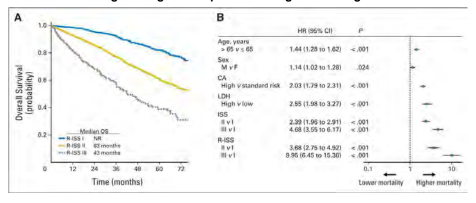






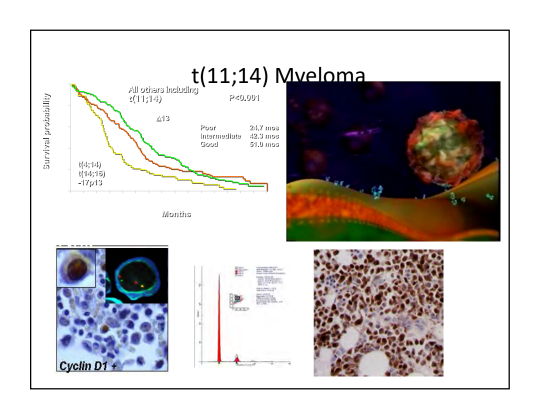
Revised ISS Model

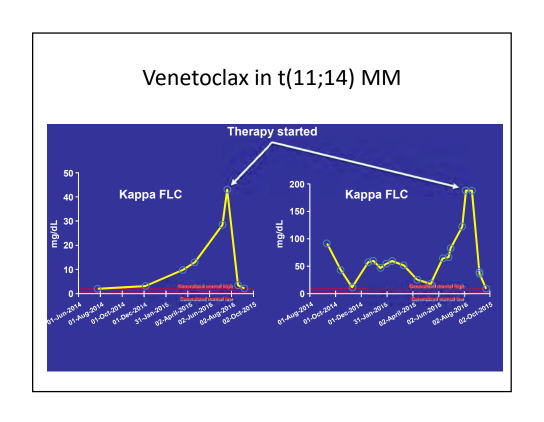
- · Revised system based on ISS (b2-microglobulin and albumin)
- · Incorporates high risk cytogenetics and elevated LDH
- · Creates model
 - · Stage 1 Stage I ISS and cytogenetics and LDH favorable
 - Stage 2 The rest
 - · Stage 3 Stage III ISS plus either high risk or high LDH

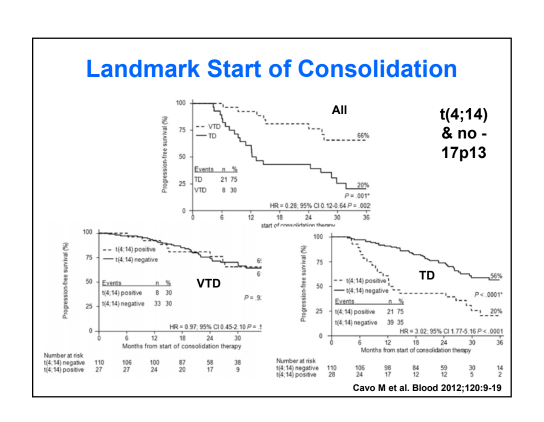


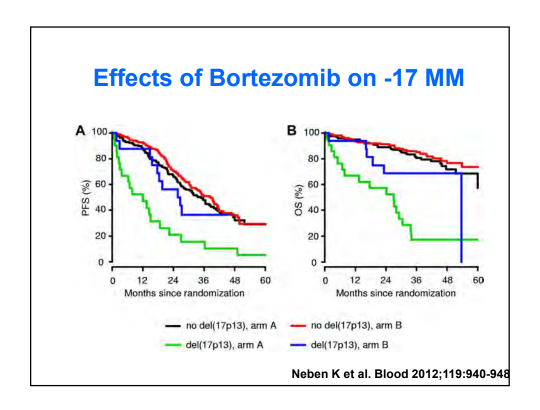
Why bother with genetics?

- Important for counseling
- Important for therapy selection
- Becoming predictive
- Don't waste money on
 - Cytogenetics
 - Flow cytometry

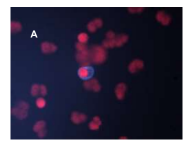




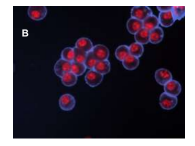




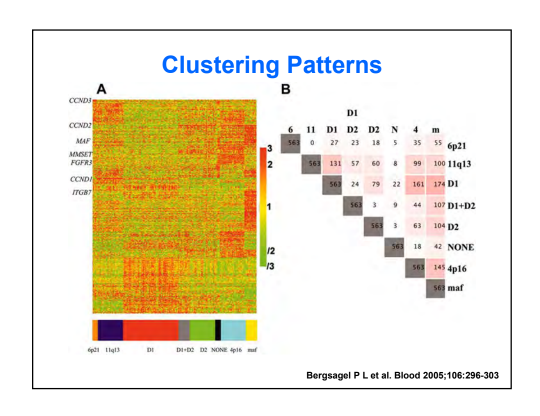
For Genomic Studies Need to Sort Cells

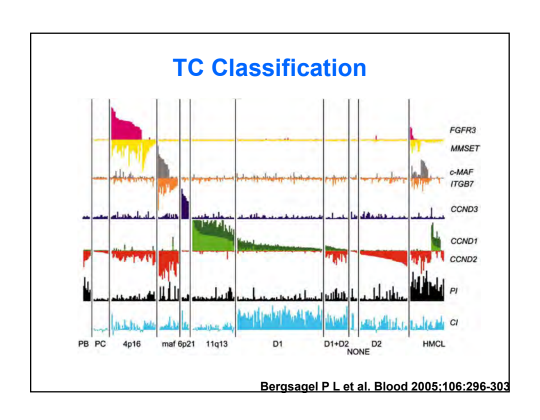


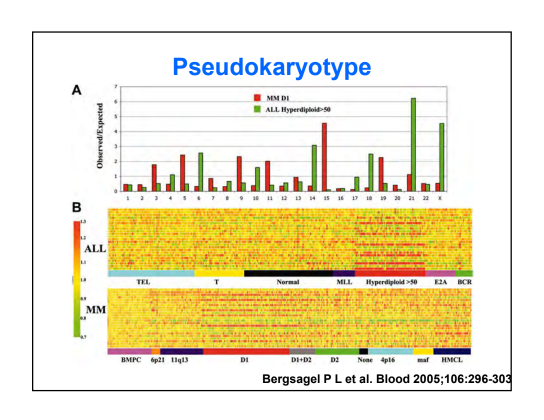
A: An ACK (ammonium chloride) lysed bone marrow from a patient with 6% lambda (AMCA) positive myeloma cells. The blue cytoplasm is AMCA conjugated Goat anti human lambda light chain. The cell nuclei were counterstained with Propidium Iodide (red).

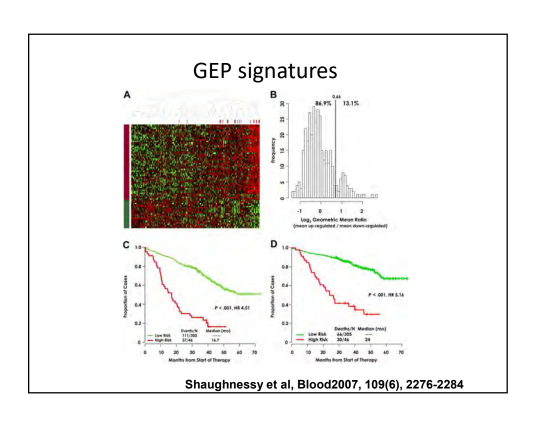


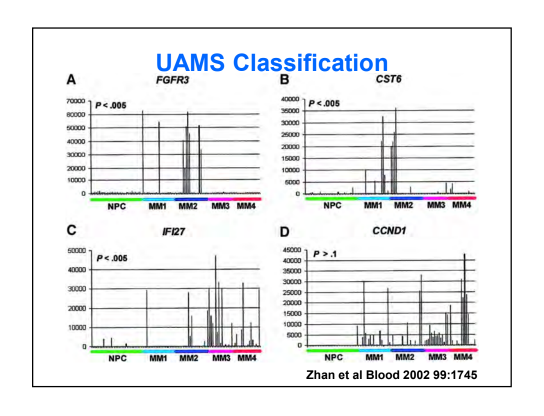
B: The same patient after CD138 bead selection using the Miltenyi autoMACS. Cells were stained with AMCA conjugated Goat anti-human lambda light chain (blue). The cell nuclei were counterstained with Propidium lodide (red). The sample was determined to be >95% Lambda positive plasma cells.

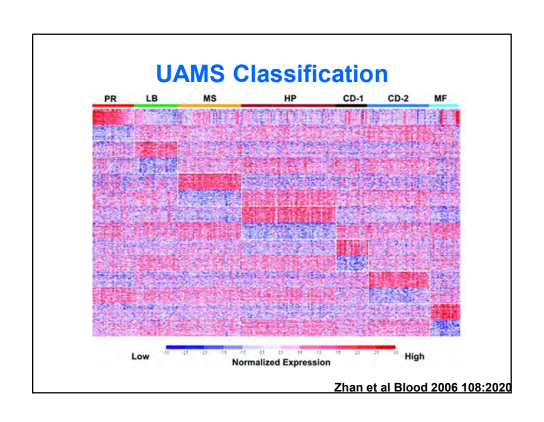


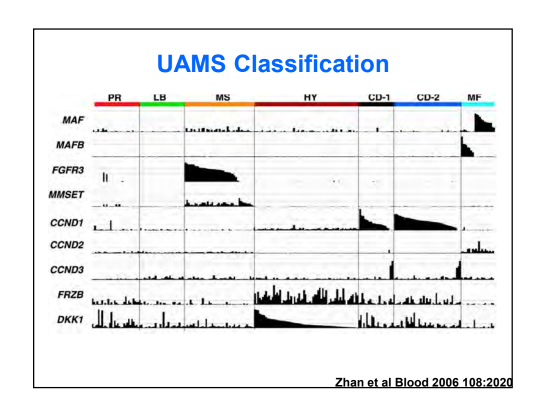


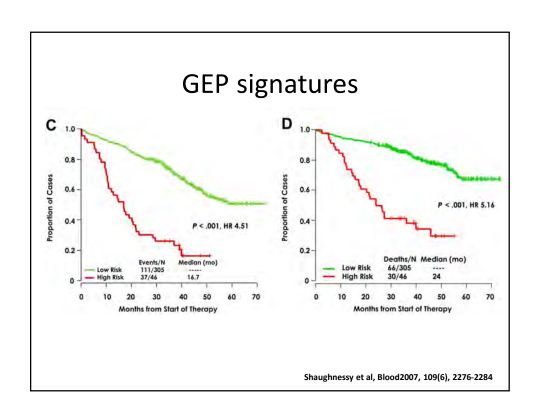


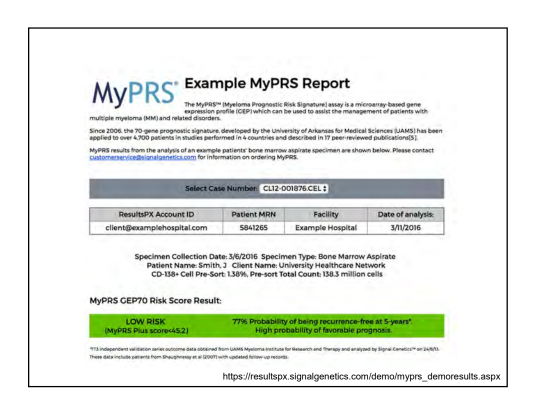


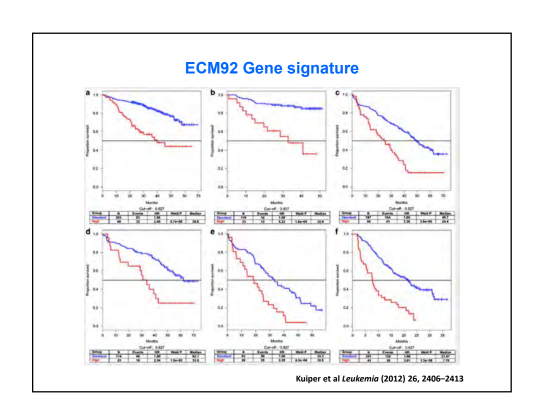


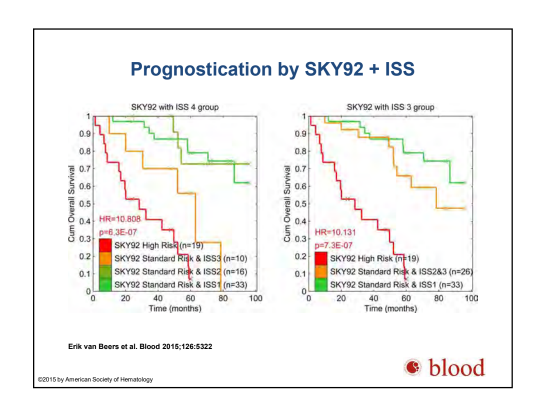


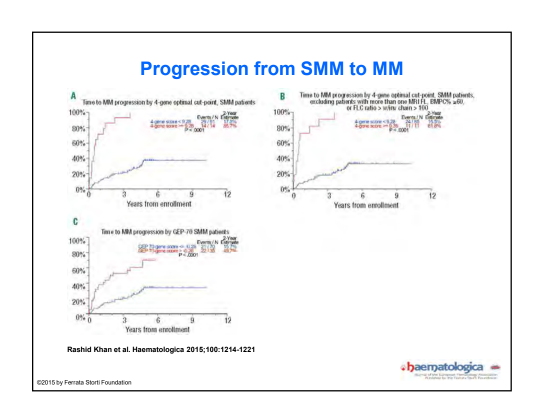












Myeloma Achilles Heel

Plasma cell biology?

Translocations?

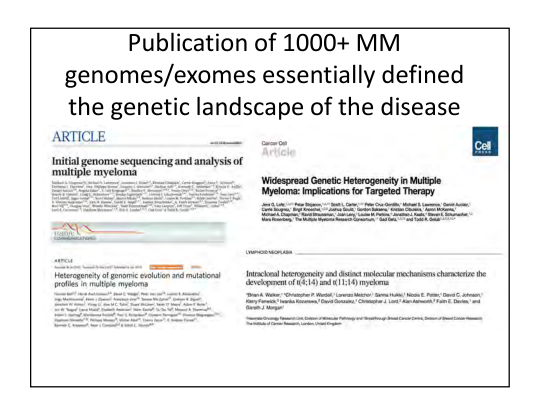
Mutations?

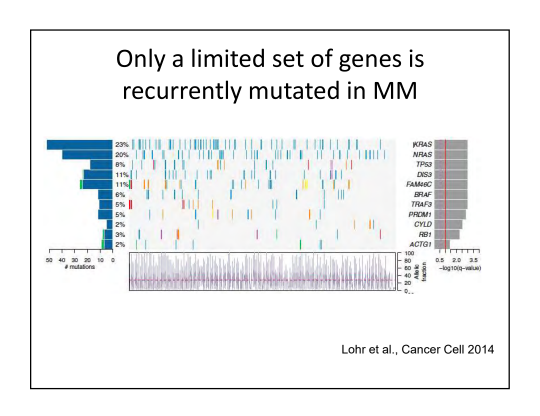
Progression events?

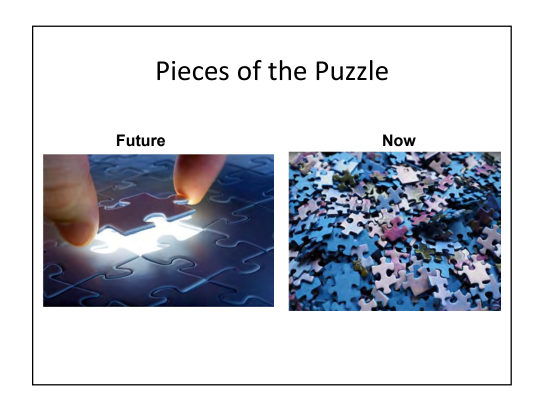
Immunology?

Microenvironment?







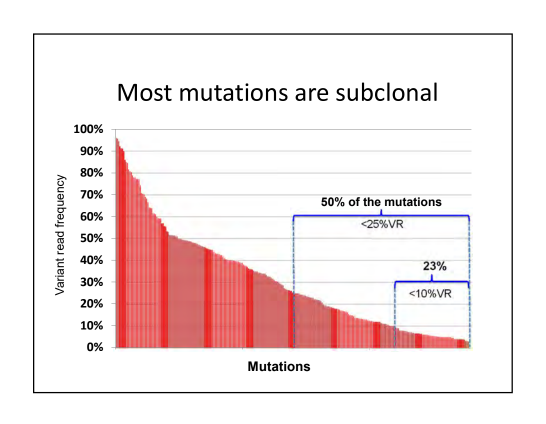


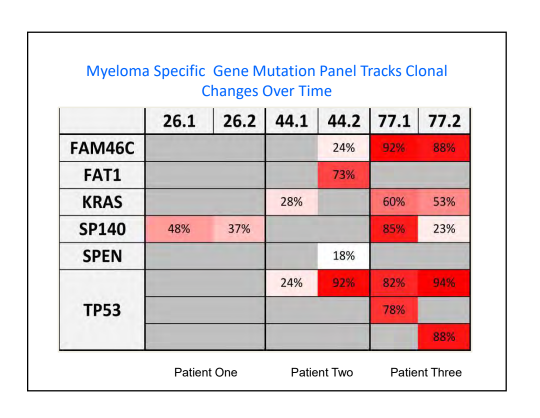
Myeloma Mutation Panel (M³P)

- Recurrently mutated putative MM genes: FAM46C, TP53, DIS3
- Actionable genes: BRAF, IDH1
- Pathways: NF-kB, MAPK, MYC
- Drug Resistance: IMiDs, Pls, glucocorticoid
- Copy-number changes
- Biallelic deletions of TSG
- Sample purity measurement: J regions IGH, IGL-K and IGL-L →88 genes, 1327 amplicons, 373 Kb

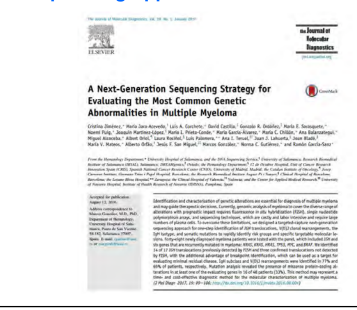
Results to Date

- > 600 tumor and germline samples average 658X
- 83% have mutation average 2 mutations/patient
- 72 untreated del17p (p53 mutation in 28%)
- Braf in 9% at diagnosis and 18% at relapse





Sequencing Approaches to Detect Genetics



Conclusions

- Genetics can help classify MM subgroups
- Genetics have powerful prognostic implications
- Combined with standard clinical factors can best stratify patients
- Clinical tests are available
- Quality of process is critical



Institute of Biomedical Research of Salamanca



University of Salamanca



Cancer Research Center

What are the optimal imaging techniques in

María-Victoria Mateos

Myeloma?

University Hospital of Salamanca University of Salamanca Spain

What is the role of imaging in Myeloma?

- Precise identification of bone disease, as sign of organ damage and need to start treatment
- ➤ Identification of sites of extra-medullary disease (total body techniques)
- ➤ Differential diagnosis between localized disease (BSP) and systemic disease (MM)
- > Correct identification of sites of bone disease at risk of complications (fractures, neurological complications) (MRI gold standard)
- > Correct follow up of the patients after treatment

Zamagni E. et al, BJH 2012

What is the role of imaging in Myeloma?

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Zamagni E. et al, BJH 2012

ACTIVE MYELOMA: the CRAB CRITERIA

Myeloma-related end organ damage due to the plasma cell proliferative process

- C: Calcium levels increased
- R: Renal insufficiency
- A: Anemia
- B: Bone lesions, osteolytic or osteoporosis

X-ray was the standard of care for bone lesions detection

Panel: Revised International Myeloma Working Group diagnostic criteria for multiple myeloma and smouldering multiple myeloma

Definition of multiple myeloma

Clonal bone marrow plasma cells >10% or biopsy-proven bony or extramedullary plasmacytoma* and any one or more of the following myeloma defining events:

- · Myeloma defining events:
 - Evidence of end organ damage that can be attributed to the underlying plasma cell proliferative disorder, specifically:
 - Hypercalcaemia: serum calcium >0.25 mmol/L (>1 mg/dL) higher than the upper limit of normal or >2.75 mmol/L (>11 mg/dL)
 - Renal insufficiency: creatinine clearance <40 mL per min† or serum creatinine >177 µmol/L (>2 mg/dL)
 - Anaemia: haemoglobin value of >20 g/L below the lower limit of normal, or a haemoglobin value <100 g/L
 - Bone lesions: one or more osteolytic lesions on skeletal radiography, CT, or PFT-CT±
 - · Any one or more of the following biomarkers of malignancy:
 - Clonal bone marrow plasma cell percentage* ≥60%
 - Involved:uninvolved serum free light chain ratio ≥100
 - >1 focal lesions on MRI studies¶

Raikumar et al. Lancet Oncology 2014: 15: e538-48

International Myeloma Working Group updated criteria for the diagnosis of multiple myeloma

SVincent Rajkumar, Meletios A Dimopoulos, Antonio Palumbo, Joan Blade, Giampaolo Merlini, Maria-Victoria Mateos, Shaji kumar, Jens Hillengass. Efstathios Kastritis, Paul Richardson, Oki Landgren, Bruno Paika, Angda Disperzien, Brendan Weiss, Xavier LeLeu, Sonja Zweagman, Sagar Lonkal. Laura Rosiinol, Bena Zamagni, Sundar jajannath, Orhan Sezer, Sigurdur Y Kristinsson, Jo Caers, Saad Z Usmani, Juan José Lativerta, Haris Erik Johnsen, Meral Beksus, Michele Cava, Hartmut Goldschmidt, Evangdos Terpor, Robert A Kyle, Kenneth C Anderson, Brian G M Durie, Jeaus F San Miguel

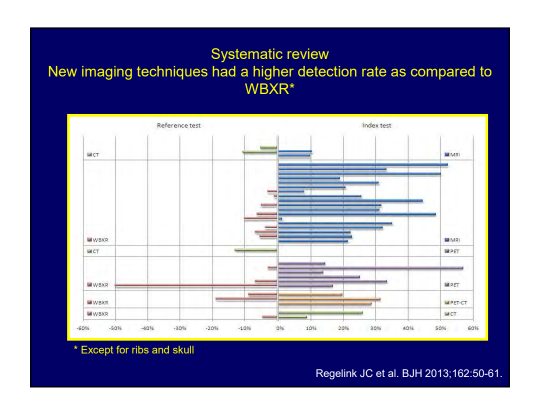
This International Myeloma Working Group consensus updates the disease definition of multiple myeloma to include validated biomarkers in addition to existing requirements of attributable CRAB features (hypercalcaemia, renal failure, anaemia, and bone lesions). These changes are based on the identification of biomarkers associated with near inevitable development of CRAB features in patients who would otherwise be regarded as having smouldering multiple myeloma. A delay in application of the label of multiple myeloma and postponement of therapy could be detrimental to these patients. In addition to this change, we clarify and update the underlying laboratory and radiographic variables that fulfil the criteria for the presence of myeloma-defining CRAB features, and the histological and monoclonal protein requirements for the disease diagnosis. Finally, we provide specific metrics that new biomarkers should meet for inclusion in the disease definition. The International Myeloma Working Group recommends the implementation of these criteria in routine practice and in future clinical trials, and recommends that future studies analyse any differences in outcome that might occur as a result of the new disease definition.

www.thelancet.com/oncology Vol 15 November 2014



International Myeloma Working Group updated Criteria for the diagnosis of Multiple Myeloma

- •Definition of myeloma bone disease (CRAB):clear evidence of one or more sites of osteolytic bone destruction (at least 5 mm or more in size) seen on CT, WBLDCT, PET/CT, regardless of weather they can be visualized on skeletal radiography or not
- •If doubt lesions on CT or PET/CT: close follow-up every 3-6 months and/or biopsy of the lesion
- •Oseoporosis per se in the absence of lytic lesions is not sufficient for CRAB



MRD supersedes CR

and could meet some of the key requirements for a surrogate endpoint such as:

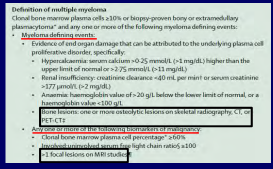
- o move the PFS of patients in remission from 3–5 years to 8–10 years
- o independence from treatment
- o predict different outcomes upon different MRD-negative rates
- o useful in all patient subgroups
- o reliable and widely available techniques, inside and outside the BM

San Miguel J. EHA 2016

ACTIVE MYELOMA: the CRAB CRITERIA

Myeloma-related end organ damage due to the plasma cell proliferative process

- C: Calcium levels increased
- R: Renal insufficiency
- A: Anemia
- B: Bone lesions, osteolytic or osteoporosis



Rajkumar V. et al., Lancet Oncology 2014

IMWG, BJH 2003

38

Is conventional radiography "the gold standard" for depicting myeloma osteolytic lesions in 2016?



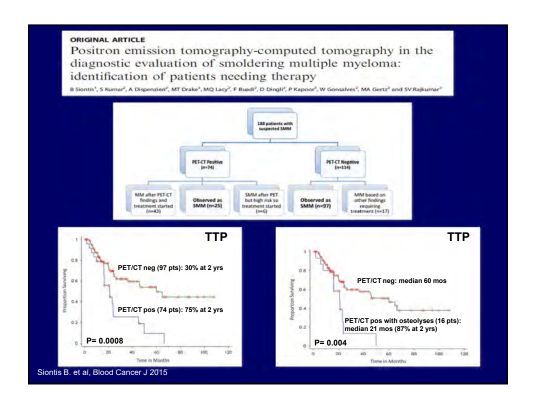


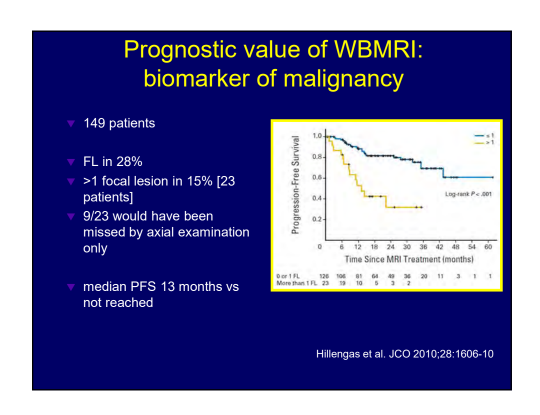


- Lytic lesions are visible only if at least 30%-50% of trabecular substance is lost
- Unable to identify small osteolytic lesions (planar technique)
- Low sensitivity in the spine
- Unable to distinguish between osteoporotic vertebral fractures and MM related ones
- It cannot be used for the assessment of response to treatment

Frequent underestimation of MM bone disease

Zamagni E. et al, BJH 2012 Pianko et al, Clin Canc Res 2014





Prognostic value of WBMRI

ORIGINAL ARTICLE

Predictive value of longitudinal whole-body magnetic resonance imaging in patients with smoldering multiple myeloma

Variable	HR	95% Cl	P-value	
1st MRI ≥2 FL	2.24	0.84-5.98	0.108	
MRI-PD	14.1	5.06-39.3	< 0.001	
M-Protein ≥20 g/l	1.05	1.01-1.09	0.022	
1st MRI ≥ 2 FL	2.90	0.45-18.6	0.260	
MRI-PD	10.4	2.57-42.0	0.001	
aPC/BMPC≥95%	6.40	1.36-30.2	0.020	

Abbreviations: aPC, aberrant plasma cells; BMPC, bone marrow plasma cells CJ, confidence intervals; FL, focal lesions; HR, hazard ratio; MRI, magnetic resonance imaging; MRI-PD, radiological progressive disease. Multivariate analysis for serum M-Protein and aPC/BMPC ≥ 95% at irritial MRI as well as radiological progressive disease (MRI-PD) for progression into symptomatic MM. P-values are derived from Wald-test in Cox proportional hazard analysis Bold and Italic numbers represent significant findings.

Newer imaging techniques: are all of them similar?

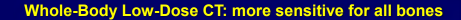
- •Morphological: assessing bone destruction
 - WB-MDCT-LDCT, CT part of PET/CT
- Functional: assessing bone marrow infiltration and disease metabolism
 - · ASSIAL MRI-WBMRI (DCE-MRI, DWI-MRI), PET/CT

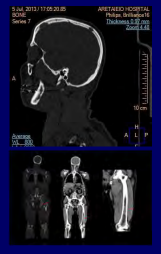
ROLE OF NEWER IMAGING TECHNIQUES

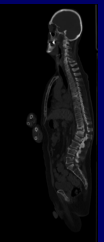
- •MORPHOLOGICAL: assessing bone destruction
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Active MM

- at diagnosis: staging and prognosis
- after treatment: evaluation of treatment response
- •Early stage/smoldering MM





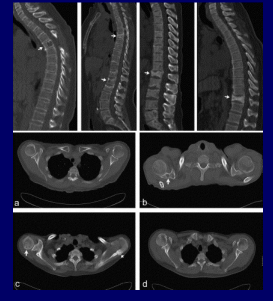






- Fast scanning time, low radiation dose (3,3-7 msV), high resolution images
- Demonstration of extra-osseus findings

WHOLE BODY LOW-DOSE MULTIDETECTOR ROW-CT (WB-LDCT)



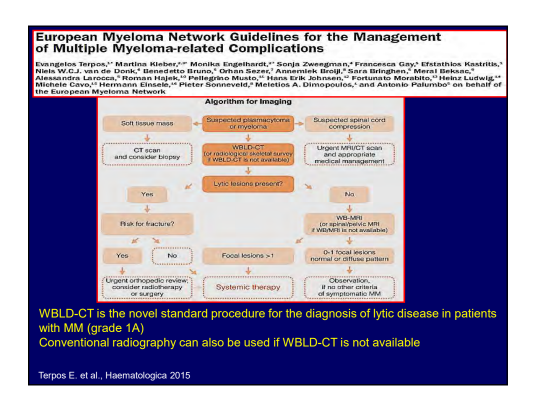
- Fast scanning time, low radiation dose (3,3-7 msV), high resolution images
- Demonstration of extraosseus findings

Shortt CP et al, Sem Musculoskel Radiology 2010 Ippolito D. et al, Eur J Radiol 2013

Wolf MB et al, Eur Journal Radiology 2014
Pianko MJ et al, Clin Canc Res 2014

Horger M., EJ Radiol, 2004 Hur J., J Comput Assist Tomogr, 2007

Study	Stud desig	•		rence est		Key find	ings	
	-	Kropil e	et al. (14)		Princewill et al. (16)			
	WBLDCT (# detected)	WBXR (# detected)	Ratio of detection (WBLDCT#/ WBXR#)	P	WBLDCT (# detected)	WBXR (# detected)	Ratio of detection (WBLDCT#/ WBXR#)	P
Total	247	120	2.06	Not reported	968	248	3.90	<0.00
skeleton Skull	7	1	7.00	Not reported	94	86	1.09	0.02
Spine	69	15	4.60	<0.001	241	49	4.92	<0.001
Thoracic cage (ribs and sternum)	60	29	2.07	<0.001	222	3	74.00	<0.00
Pelvis and flat bones	61	24	2.54	<0.001	240	36	6.67	<0.001
Long bones and extremities	46	47	0.98	Not reported	171	74	2.31	<0.001



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Rajkumar V. et al., Lancet Oncology 2014

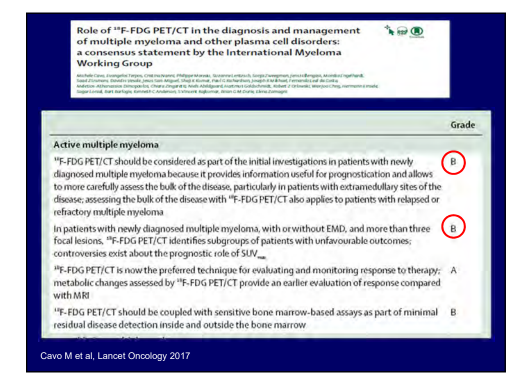
PET/CT vs WBXR OR MRI in Multiple Myeloma

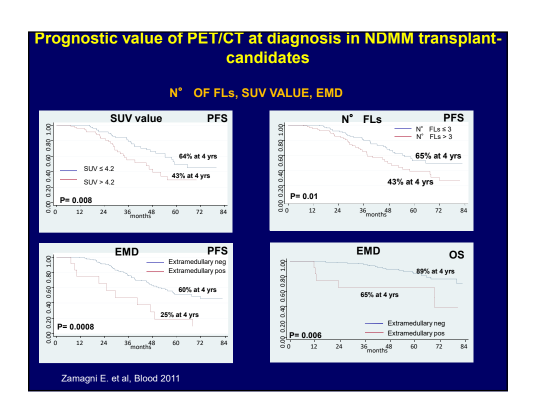
- •7 studies PET \pm CT vs WBXR: 6/7 PET showed more lytic lesions $% \left(1\right) =1$ with the exception of the skull
- •Identification of extra-medullary disease
- •CT part of PET/CT is valid for assessing bone destruction
- •PET/CT is the functional assessment for disease metabolism

PET/CT vs WBXR OR MRI in Multiple Myeloma

- •18 studies, 798 patients
- •7 studies PET \pm CT vs WBXR: 6/7 PET showed more lytic lesions $% \left(1\right) =1$ with the exception of the skull
- •5 studies PET \pm CT vs MRI spine and/or pelvis: 4/5 MRI was superior in detecting myeloma bone disease, especially in case of diffuse bone infiltration
- •1 study PET/CT vs WBMRI: concordant in 80% cases
- ·Identification of extra-medullary disease
- •CT part of PET/CT is valid for assessing bone destruction
- •PET/CT is the functional assessment for disease metabolism

Van Lammeren-Venema D et al., Cancer 2011





How to proceed in the clinic with a NDMM patient?

- Order of preference:
- PET/CT: the CT part is valid for lytic lesions detection and PET can be useful to evaluate the response to treatment
- WBLDCT: valid for lytic lesions
- X-ray: if other assessments are not available

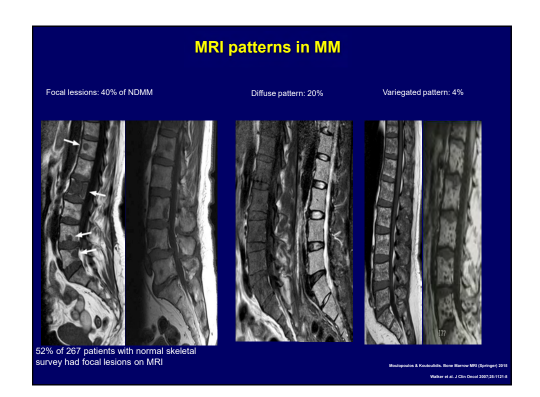
 $\bullet \textit{If doubt lesions on CT or PET/CT: close follow-up every 3-6 months and/or biopsy of the lesion \\$

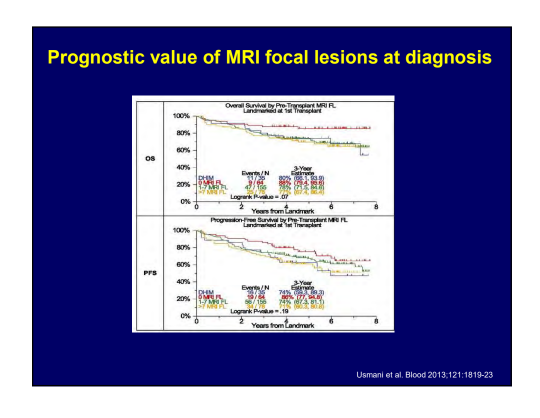
•Oseoporosis per se in the absence of lytic lesions is not sufficient for CRAB

Magnetic Resonance Imaging

- MRI is a non-invasive technique which provides detailed information about bone marrow involvement
 - > patterns: focal, diffuse, variegated, normal
 - discriminates normal vs myeloma marrow infiltration (osteoporotic vs malignant fractures)
- ➤ A WB-MRI (or MRI of the spine and pelvis) is mandatory in all patients with a presumed diagnosis of solitary plasmacytoma and in patients with smoldering MM
- ➤ It provides accurate illustration of spinal cord and/or nerve root compression, soft tissue extension, avascular necrosis

Terpos et al. J Clin Oncol 2011;29:1907-15







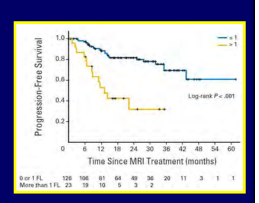
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Terpos et al. J Clin Oncol 2011;29:1907-15

Prognostic value of WBMRI in SMM: **biomarker** of malignancy

- 149 patients
- ▼ FL in 28%
- ▼ >1 focal lesion in 15% [23 patients]
- 9/23 would have been missed by axial examination only
- median PFS 13 months vs not reached



Hillengas et al. JCO 2010;28:1606-10

Imaging techniques at diagnosis in Myeloma WBLDCT PET/CT Sensitivity and specificity Sensitivity and specificity Sensitivity and specificity CT-guided biopsy, surgery, RT Optimal to assess EMD Gold standard for detection of diffuse Can depict lytic lesions (CT **BM** involvement Can depict EMD, BM Optimal for CNS imaging involvement, lytic lesions • Can assess tumor burden **PROS** Gold standard for differential diagnosis Rapid acquisition time, low and disease metabolism between osteoporotic and pathological radiation dose (3-5 mSV) • Prognostic significance of fractures Intermediate cost FLs and SUV Can depict EMD (WBMRI) · Prognostic significance of FLs Sub-optimal for diffuse BM • Imaging time (in particular axial) Sub-optimal for diffuse BM No detection of lytic lesions: not involvement CONS High cost, availability Few data/unclear prognostic enough to define end organ damage Radiation dose intermediate significance of lesion number (10 mSV) High cost , availability

How to proceed in the clinic with a NDMM patient?

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- WBLDCT: valid for lytic lesions
- X-ray: if other assessments are not available
- •If doubt lesions on CT or PET/CT: close follow-up every 3-6 months and/or biopsy of the lesion
- $\bullet \textbf{Oseoporosis per se in the absence of lytic lesions is not sufficient for CRAB } \\$
- •MRI: mandatory in solitary plasmacytoma, SMM, spinal cord compression,...

What is the role of imaging in Myeloma?

- Precise identification of bone disease, as sign of organ damage and need to start treatment
- ➤ Identification of sites of extra-medullary disease (total body techniques)
- Differential diagnosis between localized disease (BSP) and systemic disease (MM)
- Correct identification of sites of bone disease at risk of complications (fractures, neurological complications) (MRI gold standard)
- Correct follow up of the patients after treatment: Metabolic response to treatment

Zamagni E. et al, BJH 2012

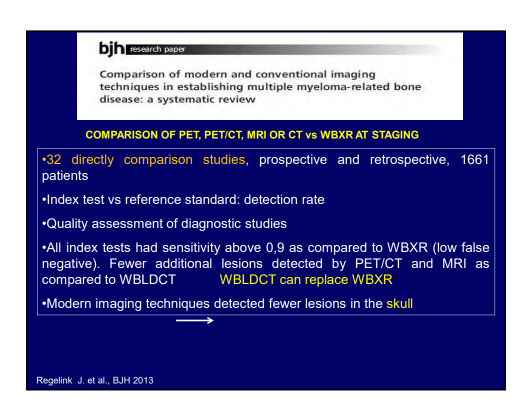
Axial MRI or DWI-WBMRI vs WBXR in Multiple Myeloma

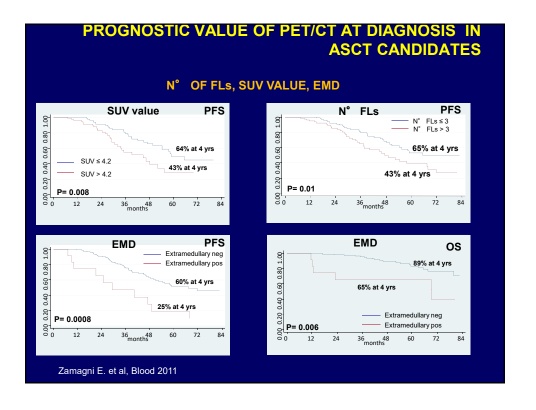
- •Detection of FLs: MRI 74%, WBXR 56%; 52% patients with normal WBXR had FLs at MRI ¹
- •WBMRI detected higher number of lesions in 37% of the patients; 18% patients WBXR negative, WBMRI positive ²
- •Clear superiority of axial MRI and WBMRI in: spine, pelvis, sternum and ribs (P < 0,001)
- •Axial MRI equally effective as PET/CT in detecting FLs (P= 0,33)³
- •Role of DWI-WBMRI in detecting the diffuse pattern^{4,5}

....but no detection of lytic lesions

²Narquin S. et al., Diagnostic and Interv Imaging 2013

^{3.} Moreau P et al, ASH 2015



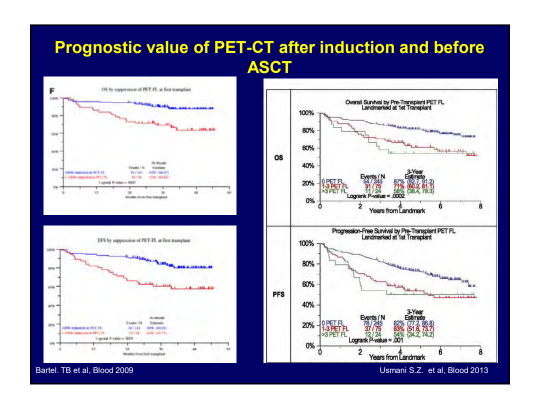


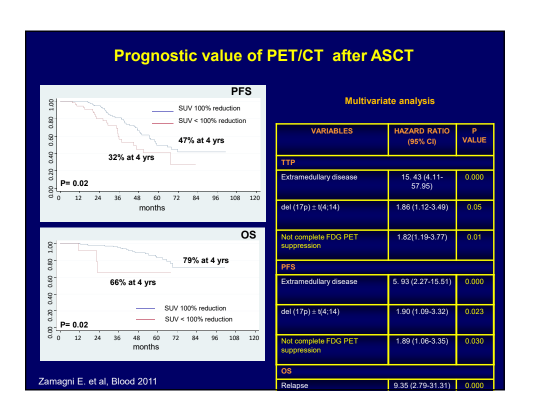
PROGNOSTIC VALUE OF PET/CT AT STAGING

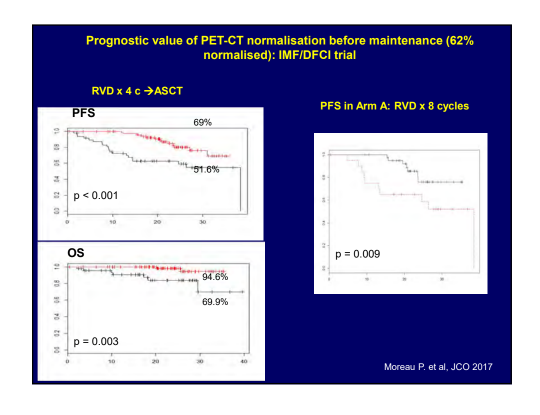
- •Several independent series of patients ASCT candidates, correlating with MRI findings, standard prognostic factors and molecular features of PCs^{1,2,3,4}
- •Small group of patients non ASCT eligible (retrospective study) ⁵
- •Series of patients pre- ALLO SCT (retrospective study) 6
- •Re-staging at relapse (retrospective studies) 7,8
- ¹ Zamagni E. et al, Blood 2011
- ² Bartel. TB et al, Blood 2009
- ³ Waheed S et al, Haematologica 2012
- ⁴ Usmani S.Z. et al, Blood 2013

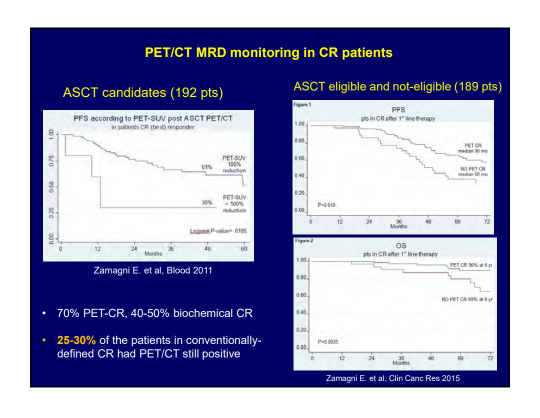
- ⁵ Zamagni E. et al, Clin Canc Res 2015
- ⁶ Patriarca F. et al, Biol BMT 2015
- ⁷ Lapa C. et al, Oncotarget 2014
- ⁸ Derlin T. et al, EJNM Mol Imag 2011

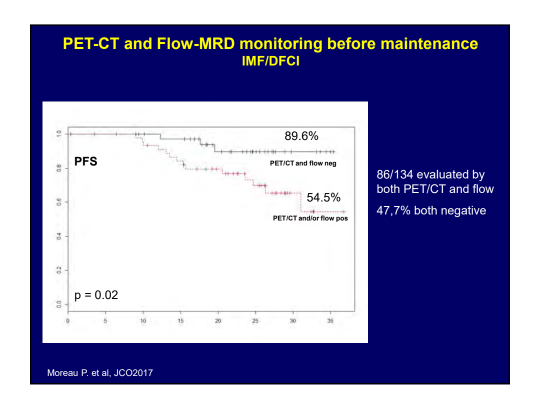
PROGNOSTIC VALUE OF MRI AT DIAGNOSIS1,2,3,4,5 TT2: CYTOGENETIC ABNORMALITIES (CA) AND MRI BOTH AFFECT OVERALL SURVIVAL OS os Overall Survival (%) Deaths / N @ Syr 60 20 Time From TT2 Enrollment (years) 48 / 191 63 / 202 95 / 218 Years from TT2 Enrollment •Correlation between diffuse MRI pattern and high-risk cytogenetics (CA) ³ •Correlation between MRI FLs and CRP, LDH, ISS 4 •Identification of a group of pts with very high-risk disease (CA+MRI) 3,4 •MRI prognostic scoring system (combining diffuse and focal infiltration pattern)⁵ ³Moulopoulos L.A. et al, AJH 2012 ¹Moulopoulos L.A. et al, Annals Oncology 2005 ²Mouolopulos L.A. et al, Leukemia 2010 ⁴Walker R. et al, JCO 2007 ⁵Mai EK. et al, Haematologica 2015

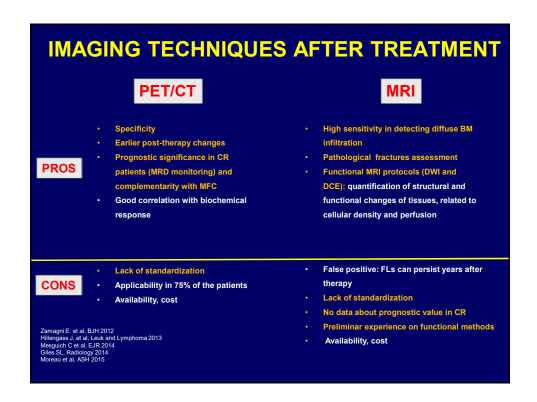




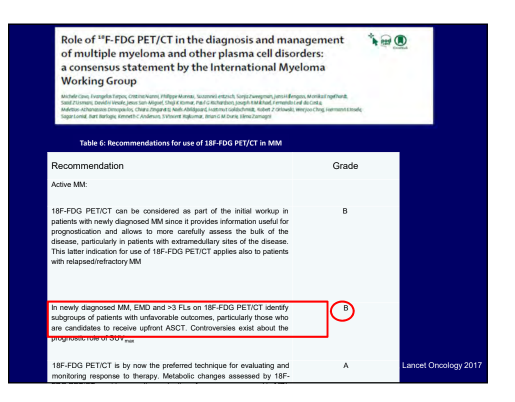








action de la	sCR (stringent complete response)	CR as defined below PLUS Normal FLC ratio 10 AND Absence of clonal cells in bone marrow biopsies by immunohistochemistry (κ/λ ratio \leq 4:1 or \geq 1:2 for κ and λ patients, respectively, after counting 2100 PCs) 7
lard IMWG Response criteria ⁶	CR (complete response)	Negative immunofixation on the serum AND urine AND ¹¹ Disappearance of any soft tissue plasmacytomas AND <5% plasma cells in bone marrow aspirates (If cellular MRD is to be performed, the first BM aspirate should be sent to MRD and morphological evaluation is not mandatory)
Respo	onse subcategory	Response criteria ¹
IMWG MRD negativity criteria (Requires CR as defined below)	Sustained MRD negative	MRD negative in the marrow (Next-generation flow or Next-generation sequencing) and by imaging as defined below, confirmed one year apart - Subsequent evaluations can be used to further specify the duration of negativity (e.g., MRD negative @ 5 years etc)
	Flow MRD- negative	Absence of phenotypically aberrant clonal plasma cells by next-generation flow cytometry ⁴ on bone marrow aspirates using the EuroFlow standard operation procedure for MRD detection in MM (or validated equivalent method) with a minimum sensitivity of 1 in 10 ⁵ nucleated cells or higher
	Sequencing MRD negative	Absence of clonal plasma cells by next generation sequencing on bone marrow aspirates in which presence of a clone is defined as less than 2 identical sequencing reads obtained after DNA sequencing of bone marrow aspirates using the Lymphosight® platform (or validated equivalent method) with a minimum sensitivity of 1 in 108 nucleated cells or higher
(Req.	Imaging+ MRD-	MRD negative as defined by Next-generation flow or Next-generation sequencing PLUS Disappearance of every area of increased tracer uptake found at baseline or a preceding PET/CT ²



PROGNOSTIC VALUE OF PET/CT AFTER TREATMENT

Study	Study design	N° pts	Treatment	PFS	os	Cox reg
Bartel Blood 2009	Р	239	ТТ3	PET-CR pre ASCT	PET-CR pre ASCT	Yes
Zamagni Blood 2011	Р	192	Thal-dex + double ASCT	PET SUV post induction and PET- CR post ASCT	PET-CR post ASCT	Yes
Usmani Blood 2013	Р	302	TT3	PET FLs (3) day +7	PET FLs (3) day +7	Yes
Dimitrakopoulou- Strauss Clin Nucl Med 2009	Р	19	СНТ	SUV during CHT	1	1
Eliott EJH 2011	R	56	Various (CHT, ASCT, novel agents)	PET-CR post therapy	1	1
Foodidarella C. et al,	Int J Mol Imaç	ging ₄ 2012	Novel agents + ASCT in 19 pts	MTV post therapy	MTV post therapy	1

PROGNOSTIC VALUE OF PET/CT AFTER TREATMENT

3 independent prospective series of patients (US, Italy, France)

- •Before ASCT (day 7 CHT, post-induction, at first ASCT)^{1,2,4}
- •After ASCT³
- •Before maintenance4

TO ASSESS MRD

- •PFS and OS difference PET pos vs neg in CR patients^{3,7} (retrospective study, 282 pts) and complementary with MFC⁴
- •No stratification of CR patients⁵ (US study, 45 pts)

¹ Bartel. TB et al, Blood 2009

² Usmani S.Z. et al, Blood 2013

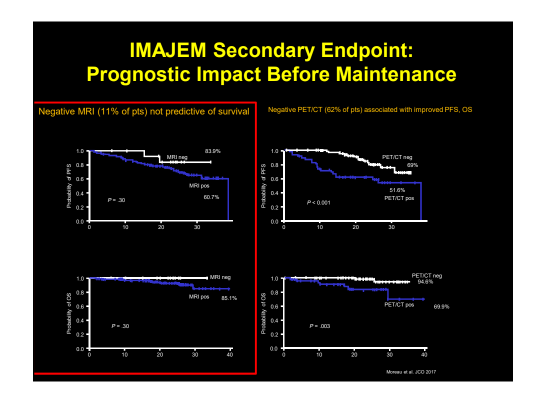
³ Zamagni E. et al, Blood 2011

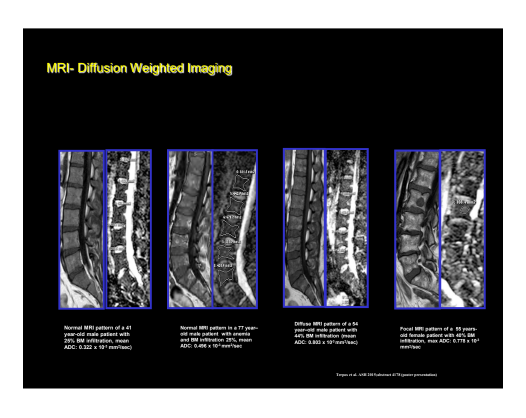
⁴Moreau P. et al, ASH 2015

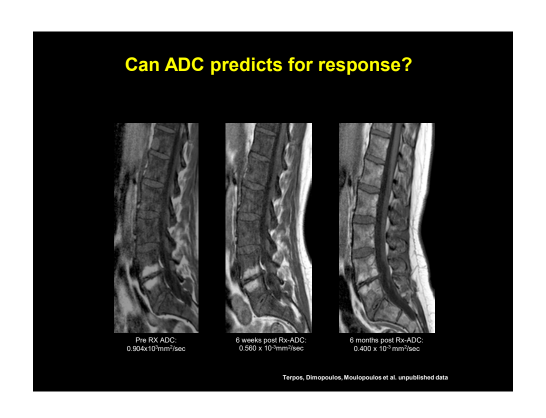
⁶ Korde N, JAMA Oncol 2015

⁷ Zamagni E. et al, Clin Canc Res 2015

Prognostic value of MRI after treatment Standard axial MRI or WBMRI •Late response after ASCT¹ (US study, axial MRI) •100 pts after ASCT² (Germany, WBMRI) **FUNCTIONAL WBMRI: DWI and DCE** Independent experiences on small series of patients (Germany, UK, France) •30 pts after CHT or ASCT: good correlation with clinical response³ (France, DCE) •26 pts after ASCT: feasibility of the technique⁴ (UK, DWI) •27 pts after treatment: correlation of DCE and DWI MRI and with clinical response •iTIMM prospective trial DWI-WBMRI vs PET/CT in newly diagnosed MM ⁴ Giles SL. et al, Radiology 2014 ¹ Walker R. et al, JCO 2007 ⁵ Bourillon C et al, Radiology 2015 ² Hillengass J. et al, Haematologica, 2012 ⁶ Dutoit JC et al, Eur J Radiol 2016 ³ Lin C et al, Radiology 2010







OPEN ISSUES

- •Quality of many studies hampered by a poor description of selection and execution criteria
- •Major inconsistency in methodology between studies
- •Need to define standardized criteria for imaging definitions and positivity cut-off

Zamagni E. et al, BJH 2012 Regelink JC et al, BJH 2013 Pianko MJ et al, Clin Canc Res 2014 Mesguich C et al, EJR 2014

IMAGING SUB-STUDY OF EMN-02

- PET/CT performed:
 - Baseline
 - After induction treatment (within 10 days)

Table-1: Five-point Deauville Criteria.

- Score 1 No uptake
- Score 2 Uptake ≤ mediastinum
- Score 3 Uptake \geq mediastinum < liver
- Score 4 Uptake moderately increased above liver at any site
- Score 5 Markedly increased uptake at any site including new sites of disease

Zamagni E. et al, ASH 2016

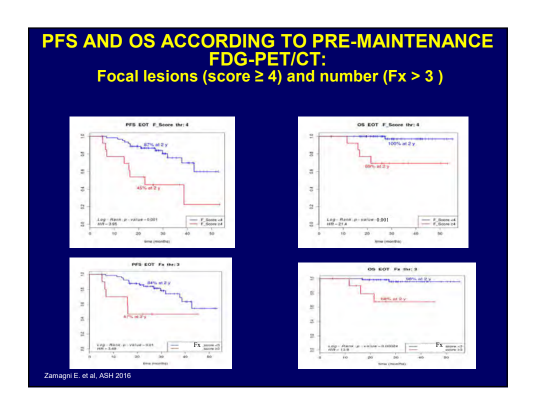
^{*} Defined as max score between Fs, BMs and EMs



- Definition of criteria for FDG-PET/CT interpretation:
 - Descriptive criteria, scored with the 5-point scale Deauville criteria for lymphomas
 - Positivity cut-off to be defined a posteriori
 - IMPeTUs vs prognosis: simplification of descriptive criteria and correlation with outcomes, according to prognostic evaluation of each parameter

Lesion type	Site	Number of lesions (x)	Grading
Diffuse	Bone marrow ^a		Deauville five-point scale
Focal (F)	Skull (S)	x = 1 (no lesions)	Deauville five-point scale
	Spine (SP)	x = 2 (1 to 3 lesions)	
	Extraspinal (ExP)	x = 3 (4 to 10 lesions)	
		x = 4 (>10 lesions)	
Lytic (L)		x = 1 (no lesions)	
		x = 2 (1 to 3 lesions)	
		x = 3 (4 to 10 lesions)	
		x = 4 (>10 lesions)	
Fracture (Fr)	At least one		
Paramedullary (PM)	At least one		
Extramedullary (EM)	At least one	N/EN (nodal/extranodal)h	Deauville five-point scale

	FDG-PET/CT PARAMETERS AND CONCORDANCE AMONG REVIEWERS											
		score 2	2		score 3	3		score 4*		score 5		
	ВМ	Fs	EM	ВМ	Fs	EM	ВМ	Fs	EM	ВМ	Fs	EM
Staging	-0.01	0.51	0.05	0.32	0.52	0.05	0.50	0.53	0.51	0.34	0.50	0.00
Post Ind	-0.04	0.46	0.34	0.24	0.46	0.24	0.38	0.45	0.22	1.00	0.38	0.00
Kripp EO1	endoi -0.03	ff's a 0.51	l <mark>pha c</mark> 0.42	o <mark>effici</mark> 0.22	ent (> 0.59	0.5 as 0.42	refere 0.41	ence) 0.50	0.49	1.00	0.31	0.45
					2	ı		1			3	
* Defined a	as > liver	uptake (S	SUVmax 3.	5)						Zamagni	E. et al, AS	SH 2016



	TE ANALYSIS			
. 50 . 2 .	, o i i , a o a a a			
	HB	lower 95	upper .95	n-velue
BM Score thr:2	1.00	1.00	1.00	0,51
BM Score thr:3	1.33	0.56	3.13	0.52
BM Score thr:4	2,17	0,49	9,61	0,31
BM_Score thr:5	1,00	1,00	1,00	0,62
Fx thr:2	2,31	0,96	5,51	0,06
Fx thr:3	3,49	1,26	9,64	0,02
Fx thr:4	3,35	0,75	14,88	0,11
F_Score thr:2	4,03	1,62	10,02	0,00
F_Score thr:3	3,65	1,51	8,85	0,00
F_Score thr:4	3,95	1,63	9,55	0,00
F_Score thr:5	24,16	2,51	232,30	0,01
EMD	4,27	1,24	14,68	0,02
Global score thr:2 (no BM)	4,28	1,66	11,07	0,00
Global score thr:3 (no BM)	3,81	1,53	9,46	0,00
Global score thr:4 (no BM)	3,65	1,54	8,69	0,00
Global score thr:5 (no BM)	2.67	0.62	11.60	0.19

FUTURE STEPS

- •Validation of the criteria in independent series of patients (FORTE GIMEMA trial) and cross-validation with french IMAJEM study (IFM 2009 trial)
- •Definition, by combined analysis of IMPeTUs and CASSIOPET, of 3 interpretative criteria:
 - · FLs at diagnosis
 - · High BM uptake
 - PET CR

Italian Myeloma criteria for Pet

Use

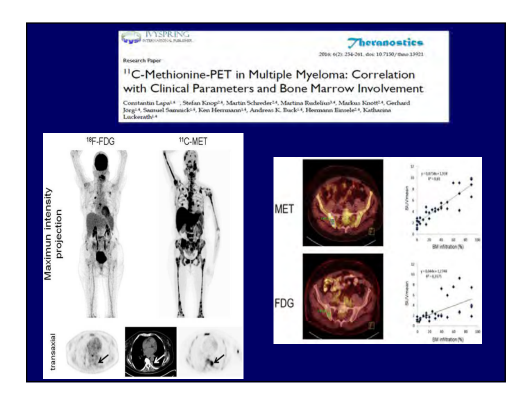
IMPeTUS

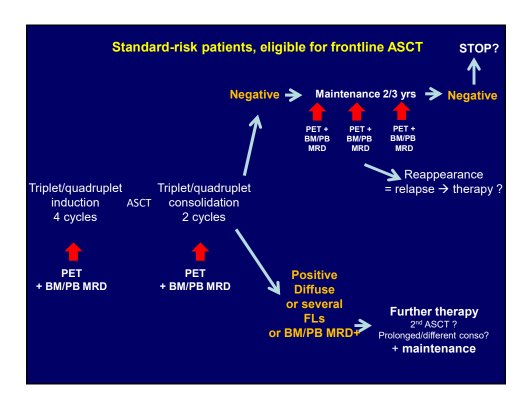
International Myeloma criteria for Pet Use

Open issues

- •How to incorporate imaging into risk-stratification at diagnosis (for both smoldering and symptomatic MM) and imaging- MRD after treatment; what is the optimal follow-up with PET?
- •What to do with persistent focal lesions after systemic therapy? What is the precise biology of persistent FLs?
- •Novel biomarkers for PET/CT (C-methionine^{1,2},C-acetate³,C-choline^{4,5,6},MM specific tracers)
- Newer techniques (PET/MRI)

¹Dankerl A. et al, Radiology 2007 ² Lapa C et al, Theranostics 2016 ³Nakamoto Y, et al, Eur J Nucl Med Mol Imaging 2013 ⁴ Lin C. et al, Eur J Nucl Med Mol Imaging 2014
 ⁵ Nanni C et al, World J Surgery Oncology 2007
 ⁶ Cassou-Mounat T, Eur J Nucl Med Mol Imaging 2016





CONCLUSION

- •Newer imaging techniques have proved reliable tools in the staging and as predictors of outcome in MM patients, both in early stage and active disease and should be used in the work-up of patients
- •PET/CT and DWI-MRI are the favorite techniques for assessing and monitoring response to therapy and are becoming complementary investigation tools for detecting minimal residual disease, going beyond the conventionally defined CR level
- •Comparative studies between PET/CT and functional MRI are warranted
- •Implementation of prospective clinical trials with newer imaging techniques will help to adress several issues, standardize the interpretation of the results and optimize the use of these promising tools



How to Perform an Appropriate Protein Screening and Decisions on Follow-up

Joan Bladé Unidad de Amiloidosis y Mieloma ICMHO, Hospital Clínic, Barcelona

Santiago de Chile, 11 de Agosto, 2017

Concept

 Clonal proliferation of mature B lymphocytes (plasma cells and/or lymphoplasmocytoid cells) resulting in a monoclonal production of an homogenous immunoglobulin (M component).

Classification

- 1. Multiple myeloma (MM) and variants
 - Smoldering myeloma
 - Plasma cell leukemia
 - Osteosclerotic myeloma (POEMS)
 - Non-secretory myeloma
 - Solitary plasmacytoma (bone or extramedullary)
- 2. Waldenström's Macroglobulinemia (WM)
- 3. Immunoglobulin light chain amyloidosis or primary amyloidosis (AL)
- 4. MG of undetermined significance (MGUS)

Incidence

- 1. MM \rightarrow 4-5 new cases/100.000 person
- 2. WM \rightarrow 0.5 new cases/100.000 person
- 3. AL \rightarrow 1 new case/100.000 person
- 4. MGUS
 - Patients >50 years-old → 3%
 - Patients >70 years-old → 5%

MM. Diagnostic

	MGUS	SMM	MM
BMPC (%)	<10 and	≥10 and/or	≥10*
M-protein (g/L)	<30	≥ 30	Any
Symptoms	No	No	Yes**

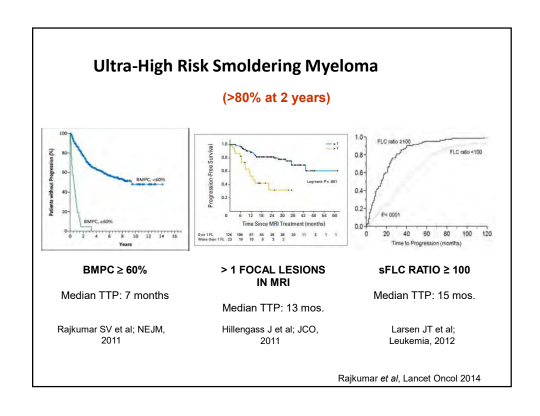
^{*}Clonal

Indicators of Increasing Disease and/or Endorgan Dysfunction MM-related (CRAB)

- HyperCalcemia (> 11.5 mg/dL)
- Renal failure (↑ serum creatinine by ≥ 2 mg/dL)
- Anemia (↓ Hb by > 2 g/dL or < 10 g/dL)
- Increase (> 50% and at least 1 cm) in size of existing Bone lesions or plasmacytomas
- Other: hyperviscosity, development of new soft tissue plasmacytomas or bone lesions

^{**}Hypercalcemia, renal failure, anemia, bone lytic lesions, recurrent bacterial infections and/or extramedullary plasmocytomas

^{*}Rajkumar et al, Blood 2011; 117: 4691-5.



Multiple Myeloma Workup

- 1. Complete blood count and differential; peripheral blood smear
- 2. Chemistry screen, including creatinine, calcium, LDH, beta2-microglobulin
- 3. <u>Serum protein electrophoresis and immunofixation</u>
- 3. <u>Serum immunoglobulins (nephelometric quantification)</u>
- 4. Measurement of serum free light chain (FLC)
- 5. <u>24-hour urine collection for electrophoresis and immunofixation</u>
- 6. Bone marrow aspirate: morphology, immunophenotype and cytogenetics by FISH (13q, t(11;14); t(4;14); t(14;16); 17p)
- 7. Radiologic skeletal survey
- 10. CT and/or MRI if clinically needed
- 11. PET/CT in patients with suspected extramedullary disease

IMWG definition of measurable disease and recommended measurements

Definitions of measurable disease

Response criteria to all categories of response except CR are aplicable only to patients who have "measurable" disease defined by at least one of the following measurements:

- Serum M-protein ≥ 10 g/L
- Urine M-protein ≥ 200 mg/24h
- Involved FLC level ≥ 100 mg/L plus an abnormal FLC ratio

Measurement of the M-protein

- Serum M-protein: quantitated using densitometry on SPEP, unless than SPEP is unrelaible, which should be explicitly reported
- Urine M-protein: quantitated using 24h-UPEP only
- Patients with "measurable disease" should be followed monthly by both SPEP and UPEP for response assessment while on therapy

IMWG Uniform Response Criteria

sCR	CR as defined below plus normal FLC ratio plus absence of bone marrow clonal plasma cells
CR	Negative IF on serum and urine, disappearance of any soft-tissue plasmacytomas and <5% BMPC
VGPR	≥ 90% serum M-protein decrease and urine M-protein <100 mg/24h
PR	≥50% serum M-protein reduction plus ≥90% urine M-protein decrease or to <200 mg/24hrs plus ≥50% reduction in the size of soft-tissue plasmacytomas

IMGW criteria for disease progression

Increase of ≥25% from lowest level in one or more of the following*:

- Serum M-protein (the absolute increase must be ≥ 5 g/L)
- Urine M-protein (the absolute increase must be ≥200 mg/24hrs)
- Bone marrow plasma cell percentage (the absolute increase must be $\geq 10\%)$
- Development of new bone lesions or soft-tissue plasmacytomas or definite increase in the size of existing bone lesions or soft-tissue plasmacytomas
- Development of hypercalcemia (corrected serum calcium >11.5 mg/dL) attributed to the plasma cell disorder

Modified from Durie et al (Leukemia 2006; 20:1467-1473)

Monitoring of Patients under Active Therapy

Induction and/or Maintenance

Before every cycle

- CBC, chemistry including serum EP and 24-hours protein urine excretion with EP
- If EP negative → serum and urine IF and, if negative, bone marrow aspirate in order to confirm CR

^{*}All categories of progression require two consecutive assessments made at any time

^{**}For progressive disease, serum M-protein increases of > 10 g/L are sufficient to define progression if the starting serum M-component was \geq 50 g/L

Follow-up in Patients with Multiple Myeloma Off-Therapy

After conventional therapy: (patients with stable response)

- First year: every 2 months
- Beyond first year: every 3 months
- Beyond 5 years: every 4 months

After HDT/SCT:

- First 6 months: every 2 months
- Two first years: every 3 months
- From 2 to 5 years: every 4 months
- Beyond 5 years: every 6 months



Patients with asymptomatic relapse or PD*

- After <u>conventional therapy</u>: every 2 months
- After <u>HDT/SCT</u>: every 3 months



Lab work-up during follow-up off therapy

- 1. Complete blood count and chemistry
- 2. Serum total protein and EF
- 3. 24-hours urine protein measurement with EF
- 4. Serum and urine immunofixation and serum FLC every 2 visits only in patients in CR
- Bone marrow aspirate and/or imaging techniques only when clinically indicated
 - Bone marrow: unexplained cytopenias (medullary progression, MDS)
 - Imaging: bone or extramedullary progression

Oligoclonal Bands

- 1. Monoclonal protein ≠ original
- 2. In patients in CR (ASCT, novel therapies)
- 3. Faint small bands in the gamma region, usually non-quantifiable
- 4. More frequent: IgG-k, IgG- λ , IgM (k or λ), k or λ light chains, rarely IgA
- 5. Frequently multiple and fluctuating
- 6. Never show a significant increase
- 7. Tipically persistent all along the CR duration and their disappearance usually precedes relapse

Is it time to implement minimal residual disease (MRD) in

multiple myeloma management?

Jesus San-Miguel
Universidad de Navarra



1st IMS workshop, Santiago de Chile 11-12 August 2017

Disclosures

Research Support/P.I. NA

Employee NA

Consultant NA

Major Stockholder NA

Speakers Bureau NA

Honoraria NA

Scientific Advisory Board NA

What should be the goal of treatment in MM?

To search for an appropriate balance between treatment efficacy, toxicity & costs

- · To eradicate the all tumor cells
- To achieve and maintain the best possible response
- MGUS signature: <10% of patients may achieve a functional cure with persistent clonal cells
 - Persistent MGUS-like clones after therapy ^{1,2}
 - Immune surveillance of residual clones 3,4



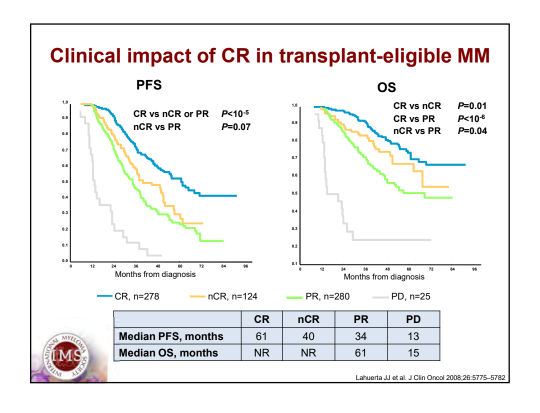
Zhan F, et al. Blood. 2007;109:1692-700
 Paiva B, et al. Leukemia. 2013;27:2056-81
 Pessoa de Magalhães RJ, et al. Haematologica. 2013;98:79-86
 Bryant C, et al. Blood Cancer J. 2013;3:e148

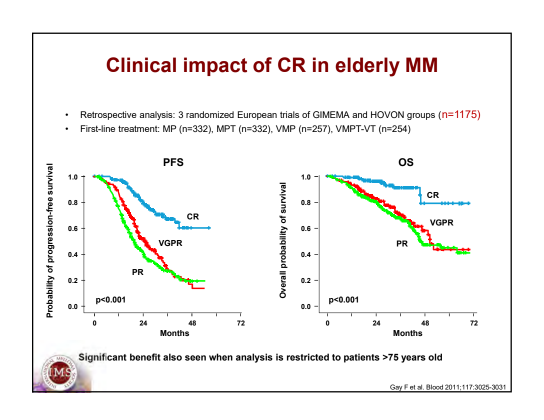
Evidence supporting this statement

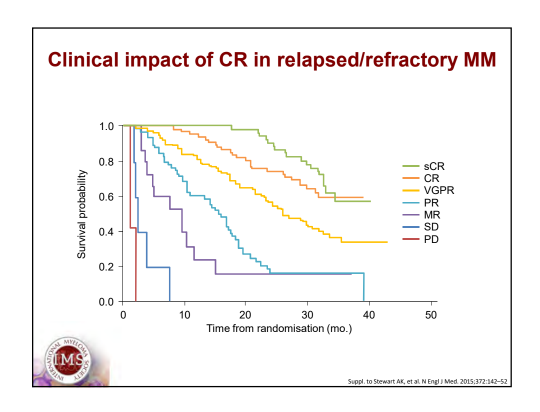
CR a surrogate marker for survival

The better the quality of the response the longer the survival









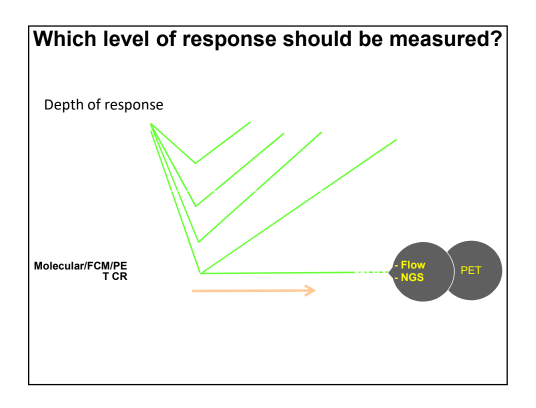
CR criteria in MM are suboptimal

➤ Negative Immunofixation & < 5% PC in BM

More sensitive techniques are needed...

MRD Techniques

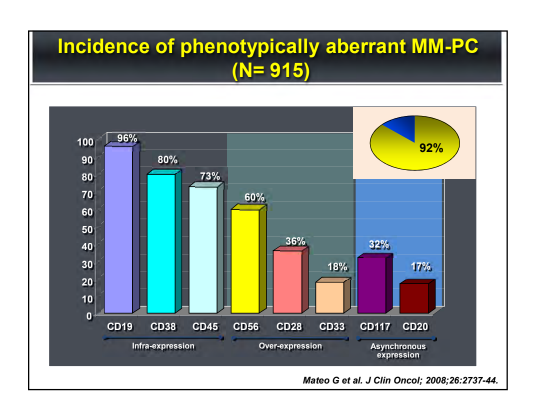


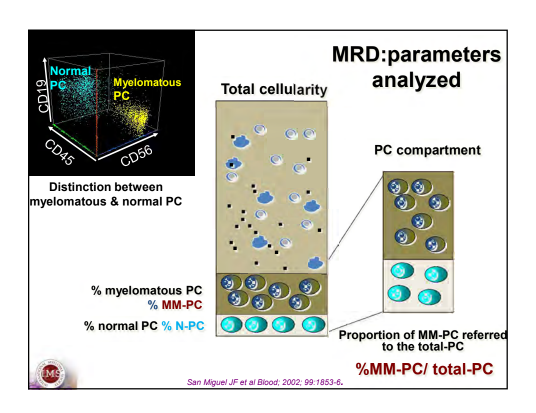


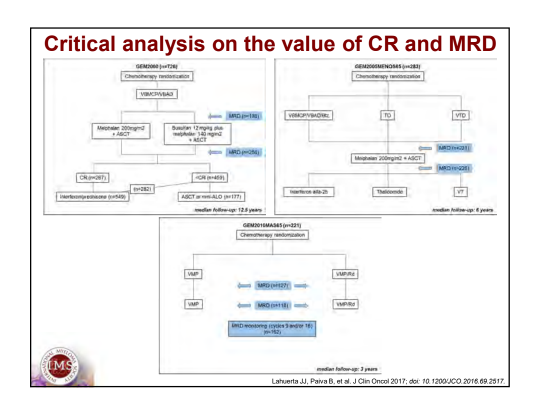
More sensitive techniques are needed.....
Multiparametric Flow Cytometry

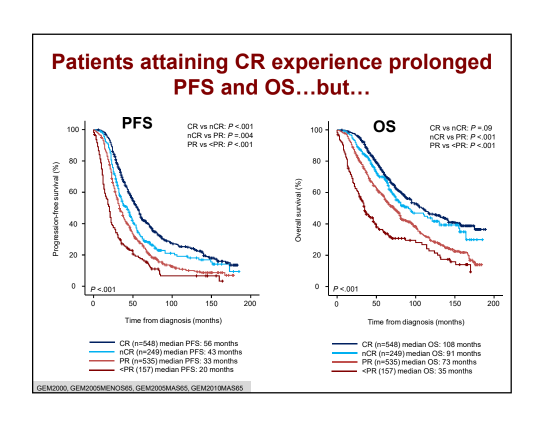
Critical analysis on the value of MRD versus CR

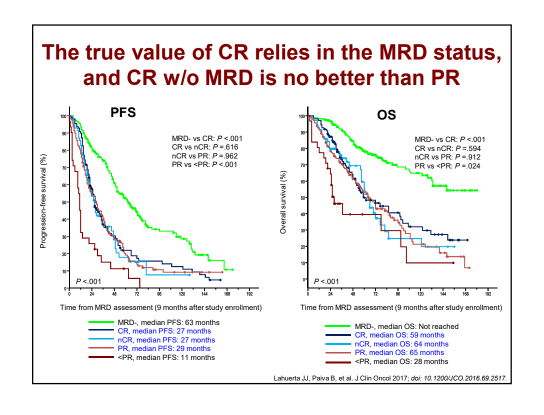


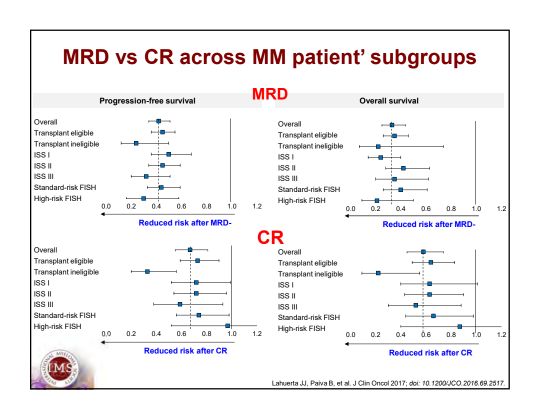




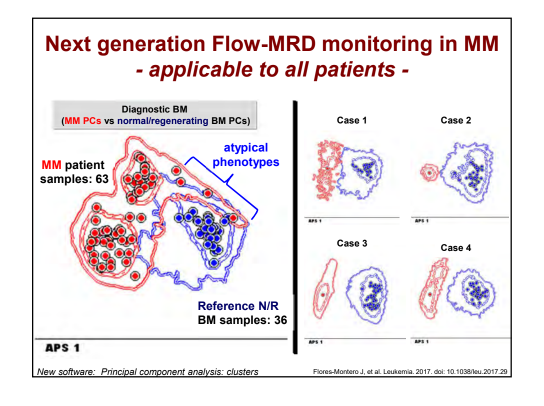


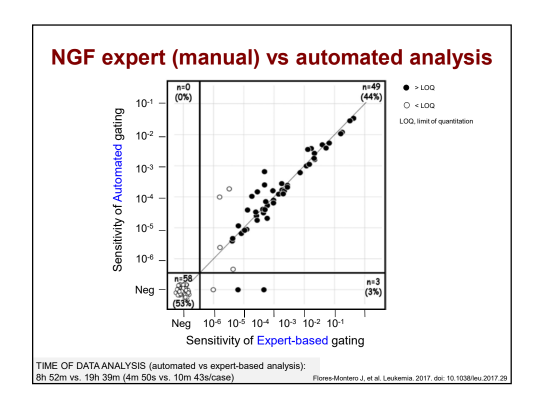


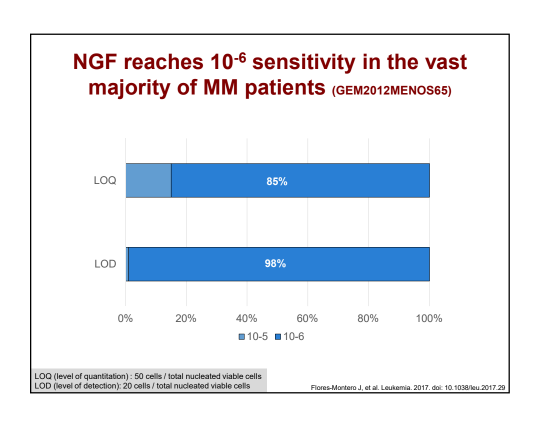


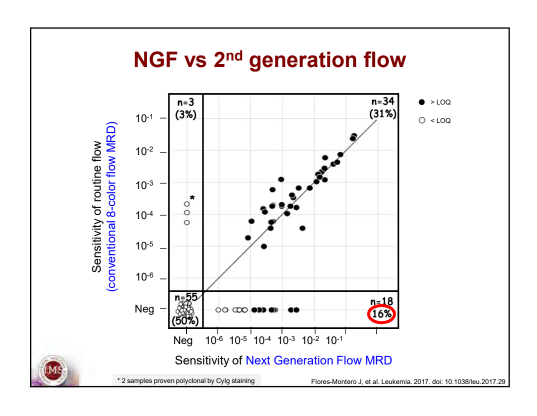


Next generation Flow-MRD monitoring in MM An optimized 2-tube 8color monoclonal antibody panel was constructed after five rounds of design-evaluation-and-redesign. Additionally, a bulk-lysis sample preparation procedure was established for acquisition of ≥10⁷ cells/patient, and novel software tools for automated analysis Flores-Montero J, et al. Leukemia. 2017. doi: 10.1038/leu.2017.29









Next generation Flow-MRD monitoring in MM

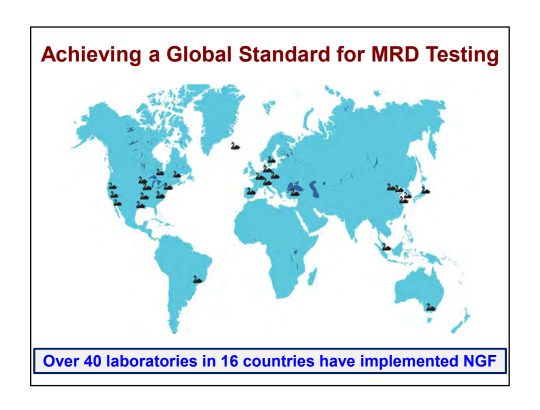
ADVANTAGES

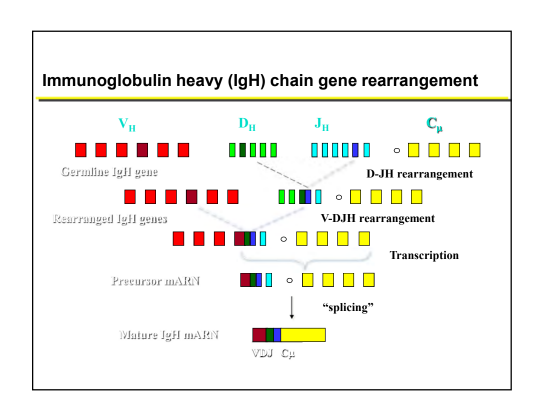
- · World-wide availability
- High applicability: 100%
- Fast for clinical decisions (<4h)
- Relatively simple
- Quantitative (% myeloma PCs)
- high-sensitivity (3x10⁻⁶)
- Assessment of non-PC BM cell compartments: sample QC
- Standardized (EuroFlow)
- · Cost (250 euro/sample)

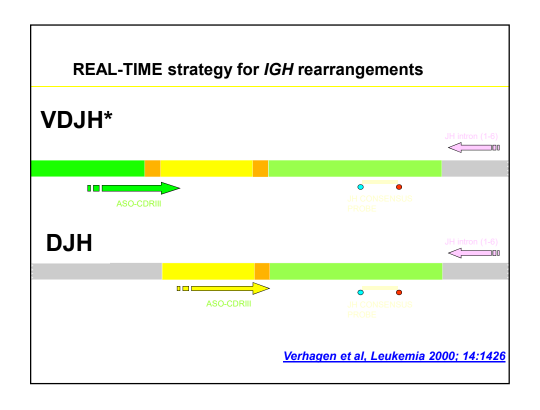
LIMITATIONS (?)

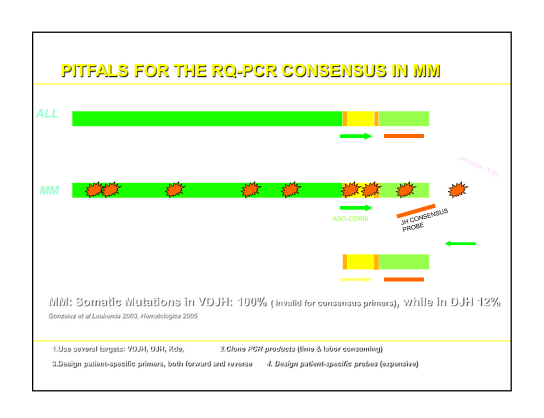
- Fresh (<48h) samples required
- Less sensitive than molecular methods
- Lack of standardization
- Heterogeneous BM infiltration or extramedullary disease





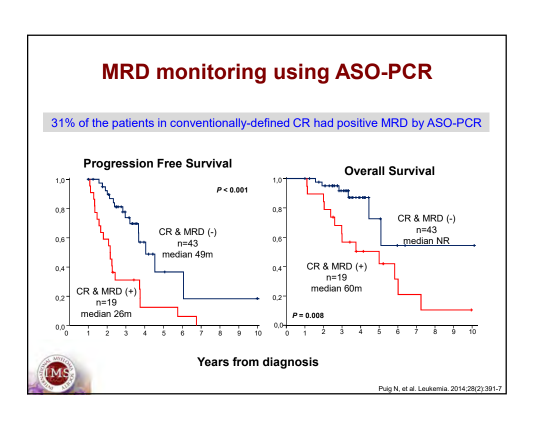






Prognostic value of MRD evaluation by PCR (Qualitative & semi-Q) in MM

Author	Context	Sensitivity	N	MRD Status	PFS	os
Corradini JCO 1999	QL ASO-PCR Auto/Allo	10 ⁻⁶	29	20 positive 9 negative	55% 78%	NR
López-Pérez et al Leukemia 2000	QL cons-PCR Auto, apheresis	10-3-10-4	27	12 positive 11 negative	20 m* 40 m	20%* 86%
Martinelli JCO 2000	QL ASO-PCR Auto/Allo	10-6	44	32 positive 12 negative	65%* 93%	NR
Corradini Blood 2003	QL ASO-PCR Allo	10 ⁻⁶	48	16 positive 19 mixed 13 negative	0% 33% 100%	NR
Ladetto et al, JCO, 2010	QL Nested-PCR VTP Post-Auto	10 ⁻⁶	39	33 positive 6 negative	66% 100%	NR
Terragna et al, ASH 2010	QL Nested-PCR VTD vs. TD post-auto	NR	67	27 positive 60 negative	NR NR	VTD: 67% TD: 52%
López-Pérez et al Leukemia 2000	Semi-QT FL-PCR Auto, apheresis	10-3-10-4	23	14 positive 13 negative	19 m* 39 m	28%* 81%
Bakkus BJH 2004	Semi-QT PCR LDM Auto	10-6	59	38 >0,015% 22 <0.015%	16 m* 64 m	NR
Martínez-Sánchez et al BJH 2008	Semi-QT FL-PCR	10 ⁻³ -10 ⁻⁴	53	25 positive 28 negative	28%* 68%	68% 86%
Puig et al Leukemia 2014	QL ASO-PCR	10 ⁻⁶	103	55 positive 48 negative	27m 54m	



MRD monitoring using ASO-PCR

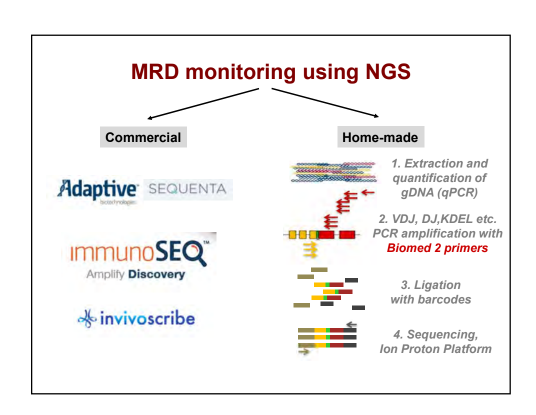


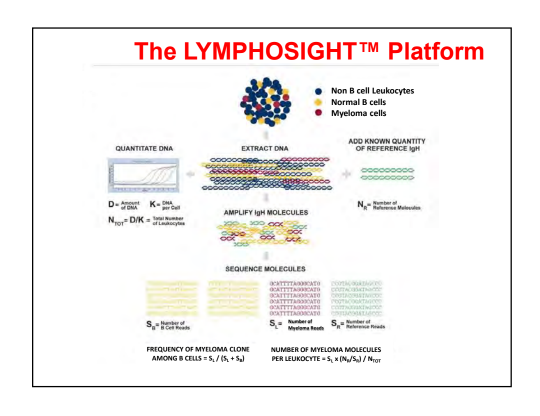


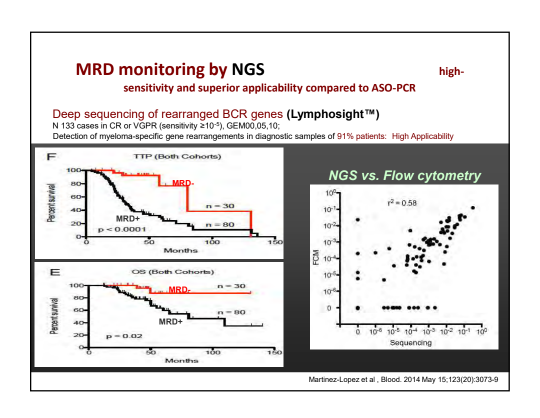
- · Highly-specific detection of clonality
- Sensitivity (10⁻⁶)
- · Detection of putative CSCs
- · Reproducibility among centers
- Does not require immediate sample processing
- Standardized (EuroMRD)

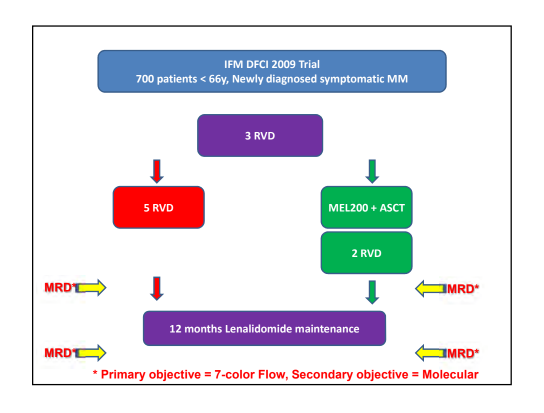
- Limited applicability (~60%)
- Limited value in patients with patchy BM infiltration and/or extramedullary disease
- Does not measure subclonal dynamics
- · No assessment of sample quality
- · Requires diagnostic sample
- · Turnaround time & time consuming
- Cost (increased by baseline sample)

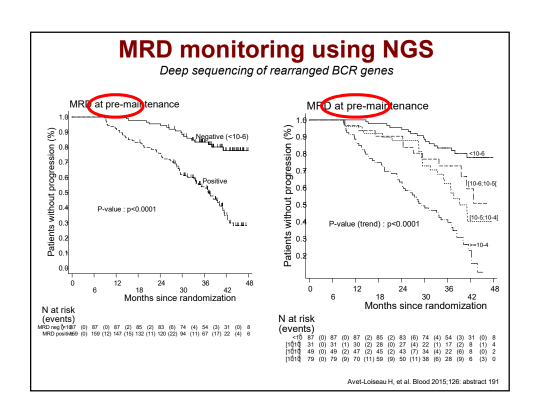


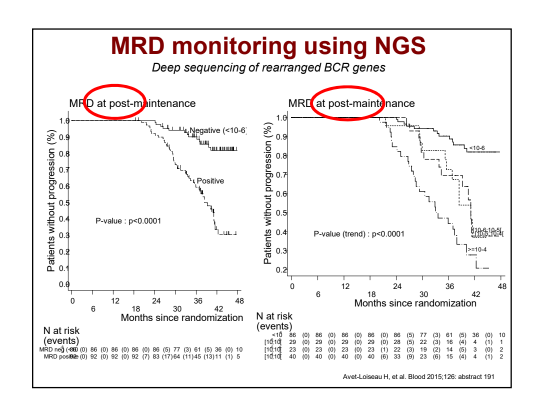


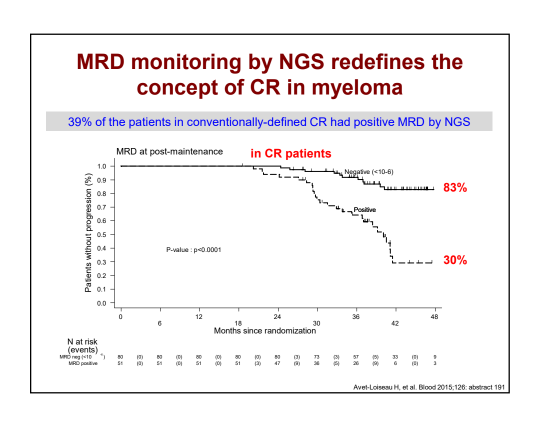


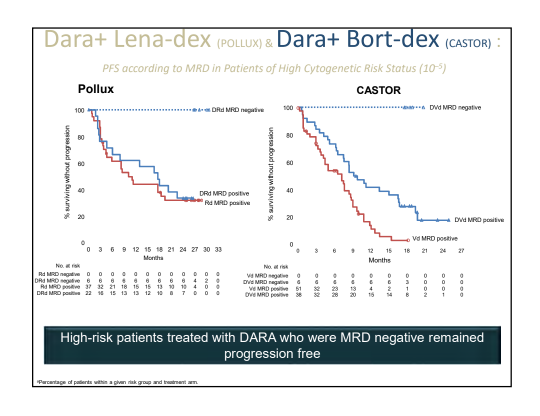


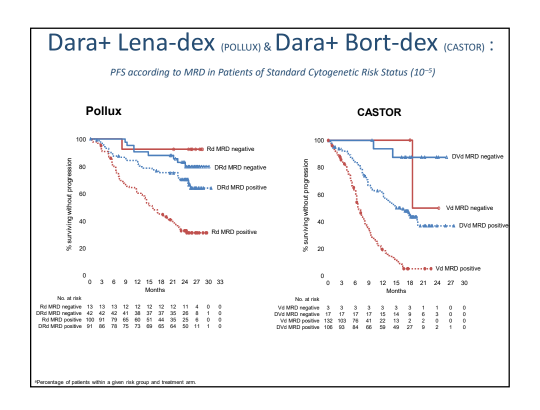












MRD supersedes **CR**

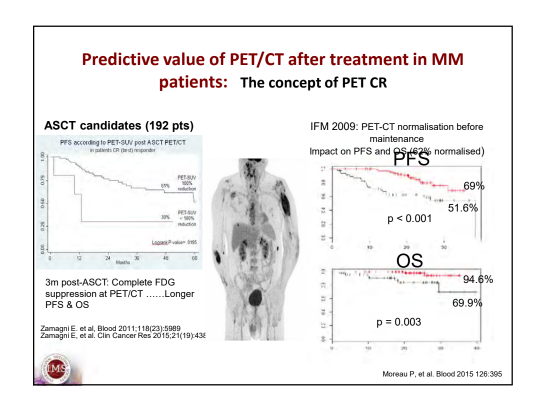
and could met some of the key requirements for a surrogate endpoint such as:

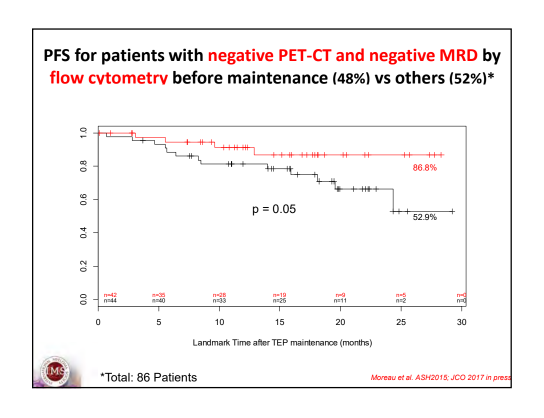
- move the PFS of patients in remission from 3-5 years to 8-10 years
- independence from treatment
- predict different outcomes upon different MRD-negative rates
- useful in all patient' subgroups
- reliable and widely available techniques



	Response subcategory Response criteria	
Sustained MRD negative in the marrow (Next-generation flow or Next-MRD generation sequencing) and by imaging as defined below, negative confirmed one year apart. Subsequent evaluations can be us to further specify the duration of negativity (e.g., MRD negatives 5 years etc.)		
ativity efined	negative	Disappearance of every area of increased tracer uptake found at baseline or a preceding PET/CT ³
RD ne	Flow MRD- negative	Absence of phenotypically aberrant clonal plasma cells by next- generation flow cytometry ⁴ on bone marrow aspirates using the EuroFlow standard operation procedure for MRD detection in MM (or validated equivalent method) with a minimum sensitivity of 1 in 10 ⁵ nucleated cells or higher
IMWG M (Requires	Sequenci ng MRD negative	Absence of clonal plasma cells by next generation sequencing on bone marrow aspirates in which presence of a clone is defined as less than 2 identical sequencing reads obtained after DNA sequencing of bone marrow aspirates using the Lymphosight® platform (or validated equivalent method) with a minimum sensitivity of 1 in 10 ⁵ nucleated cells ⁵ or higher

IMWG Criteria for MRD in Multiple Myeloma			
Response subcategory	Response criteria		
MRD negative	MRD negative in the marrow (Next-generation flow or Next-generation sequencing) and by imaging as defined below, confirmed one year apart. Subsequent evaluations can be used to further specify the duration of negativity (e.g., MRD negative @ 5 years etc)		
defined below, magning lativity criteria defined below, negative negative	MRD negative as defined below (Next-generation flow or Next-generation sequencing) PLUS Disappearance of every area of increased tracer uptake found at baseline or a preceding PET/CT ³		
MWG MRD negativity criteria Sequires CR as defined below MRD- negative Sequires CR as defined pelow MRD- negative Sequires	Absence of phenotypically aberrant clonal plasma cells by next- generation flow cytometry ⁴ on bone marrow aspirates using the EuroFlow standard operation procedure for MRD detection in MM (or validated equivalent method) with a minimum sensitivity of 1 in 10 ⁵ nucleated cells or higher		
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monitor MRD				
	NGF	ASO-PCR	NGS	PET/CT
Applicability	~100%	60-70%	~90%	~100%*
Reproducibility	High	High	Not reported	Moderate at MRD
Availability	High	Intermediate	Limited	Intermediate
Diagnostic sample	Important but not mandatory	Mandatory	Mandatory	Important but not mandatory
Time	2-3 hours	≥5 days (follow-up)	≥7 days	2-hours
Cost per sample	~350 USD	~500 USD (follow-up)	~700 USD	~2.000 USD
Sensitivity	10 ⁻⁵ – 10 ⁻⁶	10 ⁻⁵ – 10 ⁻⁶	10 ⁻⁶	High (4 mm)
Quantitative	Yes	Yes	Yes	Yes
Fresh sample	Needed	Not needed	Not needed	NA
Patchy sample	Impacts	Impacts	Impacts	No impact
Global cell characterization	Yes	No	No	No
Standardization	Ongoing (EuroFlow)	Yes, (EuroMRD)	Not reported (Adaptive) Ongoing (EuroMRD)	No

New Sensitive tests for diagnosis, prognosis and to monitor treatment efficacy

- Myeloma Diagnosis and monitoring can not remain in the Paleolithic era (....morphology and conventional Radiology)
- In MM you need to evaluate MRD inside and outside the BM
- New techniques require **standardization** (Precaution for treatment decisions...!!)
- There are **singular MM subtypes** ('rapid responders but early relapsing' or with 'MGUS profile'): in these the standard response criteria do not correlate with outcome



New Sensitive tests for diagnosis, prognosis and to monitor treatment efficacy (II)

- New techniques are expensive.... MRD follow-up study (1200 \$), PET (1300\$)......but compare this with the costs of just one additional cycle of novel drugs (sometimes no needed)......
- MRD techniques will contribute both to a better definition of response and to monitor the efficacy of intensification and maintenance therapies...may be critical to tailor treatment to avoid both under & over treatments
- But....Please do not forget Standard techniques







DISCLOSURES OF COMMERCIAL SUPPORT

Name of Company	Research support	Employee	Consultant	Stockholder	Speaker's Bureau	Advisory Board	Other
Janssen					Х	Х	
Celgene	х				Х	Х	
Amgen					Х	Х	
Takeda					х	X	

Objectives

- What is Smouldering Myeloma?
- Diagnosis of Smouldering Myeloma and differential diagnosis with other plasma cell disorders
- Why do we call it "Smouldering" Myeloma?
- What is the mechanism of transition from SMM to MM?
- What is the risk of progression to Myeloma?
- Is it possible to evaluate the individual risk of progression to Myeloma?
- What is the optimal management for Smouldering Myeloma patients?

Objectives

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Smouldering MM: diagnostic criteria					
Study	M Protein (g/dl)	Bone Marrow Plasma Cells (%)			
Kyle and Greipp, 1980	≥ 3	≥ 10			
Alexanian et al, 1988	> 2	Cor			
Wisloff et al, 1991	IgA > 1,5; IgG > 2 the G	riteria for			
Facon et al. 1995 Weber et al. 1995	ries of six patients who met the G (MM) but whose disease did not and they did not progress and they proteinuria > 1 g/24h	have an			
On the burneloma multiple myeloma	ries of sit whose disease (MM) but whose (MM) but whose disease (MM)	>10			
Ro aggressive	≥ 3	≥ 10			
IMWG, 2003∞	≥ 3	≥ 10			
*Either diagnostic criterion is acceptable, &E	oth diagnostic criteria are required, ∞Either or both diagnost	ic criteria are acceptable			

Smouldering MM: diagnostic criteria

Study	M Protein (g/dl)	Bone Marrow Plasma Cells (%)
Kyle and Greipp, 1980	≥ 3	≥ 10
Alexanian et al, 1988	> 2	
Wisloff et al, 1991	IgA > 1,5; IgG > 3	
Weber et al, 1997	> 2,5	
Facon et al, 1995	-	>15
Cesana et al, 2002*	IgA 2.1-4.9; IgG 3.6-6.9 Light chain proteinuria > 1 g/24h	>10
Rosiñol et al, 2003&	≥ 3	≥ 10
IMWG, 2003∞	≥ 3	≥ 10

*Either diagnostic criterion is acceptable &Both diagnostic criteria are required

Smouldering MM: diagnostic criteria

Study	M Protein (g/dl)	Bone Marrow Plasma Cells (%)
Kyle and Greipp, 1980	≥ 3	≥ 10
Alexanian et al, 1988	> 2	-
Wisloff et al, 1991	IgA > 1,5; IgG > 3	-
Weber et al, 1997	> 2,5	-
Facon et al, 1995	-	>15
Cesana et al, 2002*	IgA 2.1-4.9; IgG 3.6-6.9 Light chain proteinuria > 1 g/24h	>10
Rosiñol et al, 2003&	≥ 3	≥ 10
IMWG, 2003∞	≥ 3	≥ 10

*Either diagnostic criterion is acceptable &Both diagnostic criteria are required
∞Either or both diagnostic criteria are acceptable

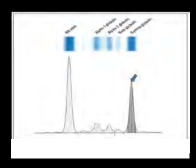
[∞]Either or both diagnostic criteria are acceptable

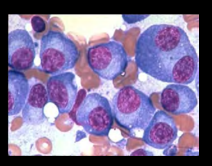
Clinical Case

- 52 years-old man
- Asymptomatic.
- Routine analysis
- Elevated total serum proteins (10.2 g/dL) with normal albumin
- Hemogram and biochemistry normal

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- Elevated total serum proteins (10.2 g/dL) with normal albumin
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Smouldering MM: diagnostic criteria

Smouldering Multiple Myeloma (SMM)

Monoclonal component

≥ 3 g/dL serum

AND/OR

Bone Marrow Plasma Cells (%) 10-60% AND

Mieloma-defining event ^a Absent

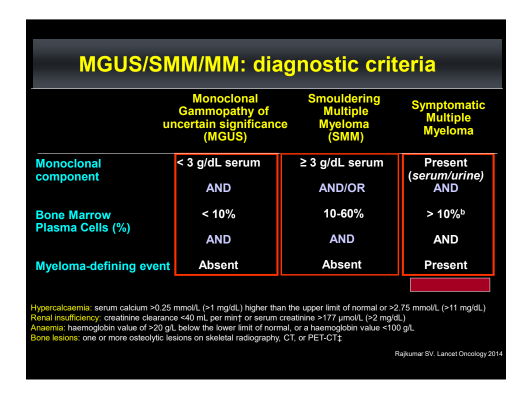
a) Mysloma Related Organ or Tissue Impairment (end organ damage) related to Plasma cell proliferative process: anemia with 2 g/dL below the normal level or 10 g/dL, or serum calcium level >10 mg/L (0.25 mmol/L), above normal or >110 mg/L (2.75 mmol/L), or lytic bone lesions or osteoporosis with compressive fractures, or renal insufficiency (creatinine >2 mg/dL or 173 mmol/L),[CRAB: Calcium increase, Renal impairment, Anemia and Bone lesion) or symptomatic hyperviscosity,, amyloidosis or recurrent bacterial infections >2 episodes in 12 m).

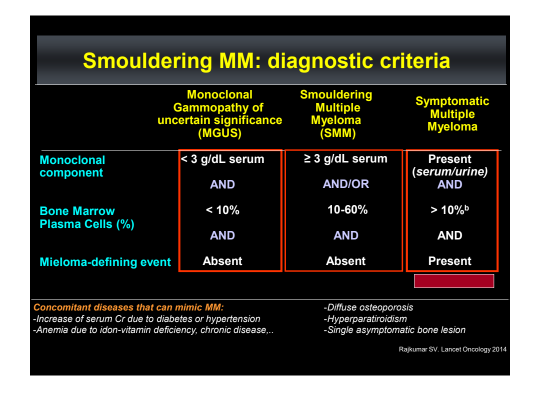
end organ damage

Rajkumar SV et al. Lancet Oncology 2014

Objectives

- What is Smouldering Myeloma?
- Diagnosis of Smouldering Myeloma and differential diagnosis with other plasma cell disorders
- Why do we call it "Smouldering" Myeloma?
- What is the mechanism of transition from SMM to MM?
- What is the risk of progression to Myeloma?
- Is it possible to evaluate the individual risk of progression to Myeloma?
- What is the optimal management for Smouldering Myeloma patients?





Recommended work up at 3 months in patients with MGUS/smouldering MM

- · Medical History and physical examination
- Hemogram
- · Creatinine and calcium values
- Protein studies
 - Total serum protein and serum electrophoresis (serum M-protein)
 - 24-h urine protein electrophoresis (urine M-protein)
 - Serum and urine immunofixation
 - Serum free light chain mesurement (FLC ratio)

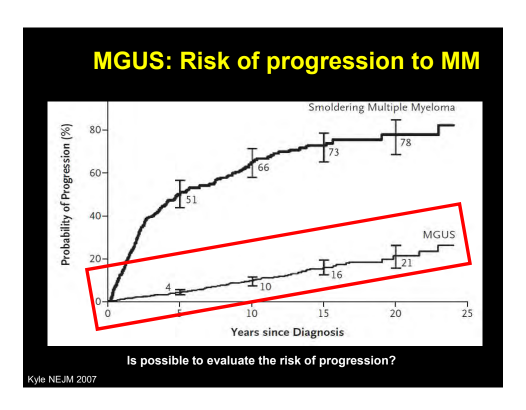
If results show stabilization of the disease, diagnosis of MGUS/SMM is confirmed

Mateos MV et al. Current hematologic malignancy reports. 2013; 8(4): 270-6

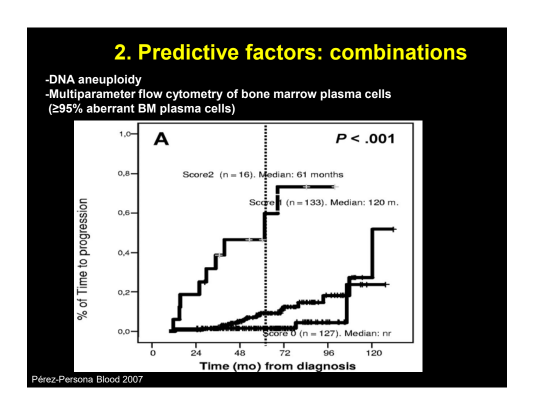
MGUS: 3 subtypes Aymptomatic; low-Asymptomatic; Symptomatic high-risk of risk of progression progression MMNon-IgM MGUS SMM IgM MGUS **SWM** WM Light-chain MGUS Light-chain MM **Idiopathic Bence** Jones proteinuria time 5.3% of population older than 70 yrs present a MGUS

Recommended work up at baseline in patients with MGUS

- · Medical History and physical examination
- Hemogram
- Creatinine and calcium values
- **Protein studies**
 - Total serum protein and serum electrophoresis (serum M-protein)
 - 24-h urine protein electrophoresis (urine M-protein)
 - Serum and urine immunofixation
 - Serum free light chain mesurement (FLC ratio)
- Bone Marrow aspirate+/- biopsy*
- Bone lesions evaluation*
- For IgG MGUS with M-protein <1.5 g/dL, bone marrow and skeletal survey are not necessary For IgM MGUS, bone marrow and CT are recommended to detect MW or NHL

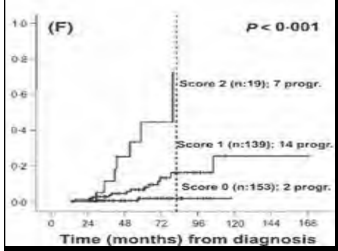


1. Predictive factors: combinations - non IgG MGUS - M-protein ≥ 15 g/L - Abnormal serum kappa/lambda FLC ratio All 3 factors abnormal Any 2 factors abnormal Any 1 factor abnormal Serum M-spike <1.5 gm/dL, IgG Subtype and normal FLC ratio



3. Predictive factors: combinations

- -Evolution of monoclonal component
- -Multiparameter flow cytometry of bone marrow plasma cells (≥95% aberrant BM plasma cells)

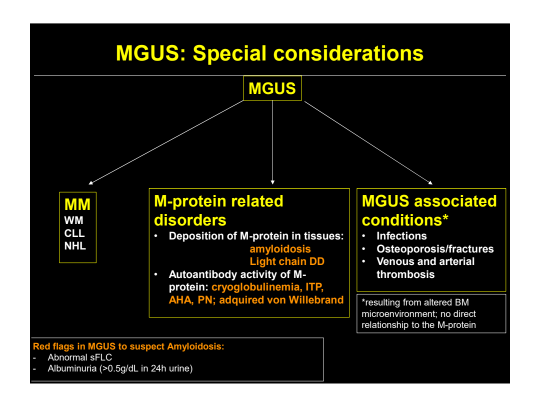


Pérez-Persona BJH 2010

MGUS: Management

- · Management should be risk-adapted
- Low risk MGUS (IgG/<1.5 g/dL/normal FLC): if stable 6 months after dx confirmation→ every 2 to 3 years
- Intermediate/high risk MGUS should be followed annually

Kristinsson et al. reported significantly better OS in patients with MM who had prior knowledge of MGUS than in those without prior knowledge (median survival, 2.8 years versus 2.1 years, respectively; P = 0.01), suggesting that earlier treatment of MM leads to improved survival



MGUS

- MGUS is one of the most common premalignant disorders
 - Malignant transformation
 - -Symptoms related to the M-protein
 - -Symtoms due to cytokines produced by plasma cell clone
 - -Symtoms related to the plasma cell clone
- · Better biomarkers for predicting progression are needed
- Treatment of high-risk patients?

MGUS prevalence

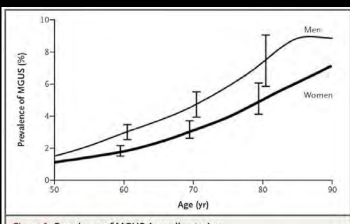


Figure 1. Prevalence of MGUS According to Age.

The I bars represent 95 percent confidence intervals. Years of age greater than 90 have been collapsed to 90 years of age.

Kyle NEJM 2006

Panel: Revised International Myeloma Working Group diagnostic criteria for multiple myeloma and smouldering multiple myeloma

Definition of multiple myeloma

Clonal bone marrow plasma cells ≥10% or biopsy-proven bony or extramedullary plasmacytoma* and any one or more of the following myeloma defining events:

- Myeloma defining events:
 - Evidence of end organ damage that can be attributed to the underlying plasma cell proliferative disorder, specifically:
 - Hypercalcaemia: serum calcium >0.25 mmol/L (>1 mg/dL) higher than the upper limit of normal or >2.75 mmol/L (>11 mg/dL)
 - Renal insufficiency: creatinine clearance <40 mL per min† or serum creatinine >177 μmol/L (>2 mg/dL)
 - Anaemia: haemoglobin value of >20 g/L below the lower limit of normal, or a haemoglobin value <100 g/L
 - Bone lesions: one or more osteolytic lesions on skeletal radiography, CT, or PET_CT+

Any one or more of the following biomarkers of malignancy:

- . Clonal bone marrow plasma cell percentage* ≥60%
- Involved:uninvolved serum free light chain ratio
 ≥100
- >1 focal lesions on MRI studies¶

Subgroup of SMM patients that require be treated

Rajkumar et al. Lancet Oncology 2014; 15: e538-48







MGUS & Smouldering Myeloma:

María-Victoria Mateos
University Hospital of Salamanca- IBSAL
Salamanca. Spain

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- Bone Marrow aspirate+/- biopsy*
- · Skeletal survey/Low-dose CT/PET-CT
- . MRI of the spine and pelvis/ Whole-body MRI*
 - *Required to identify ultra high risk SMM->MM

Mateos MV et al. Current hematologic malignancy reports. 2013; 8(4): 270-6.

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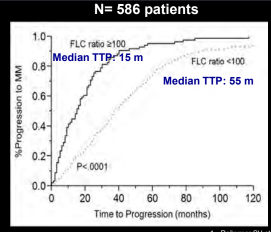
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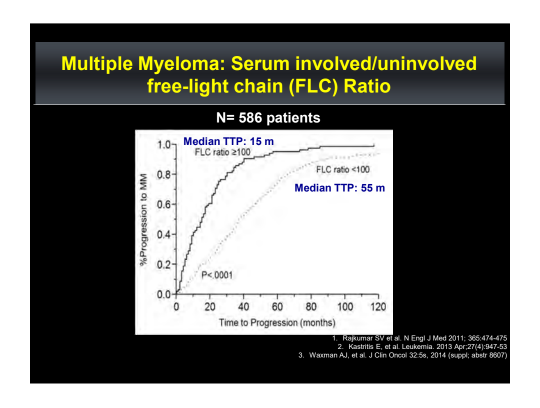
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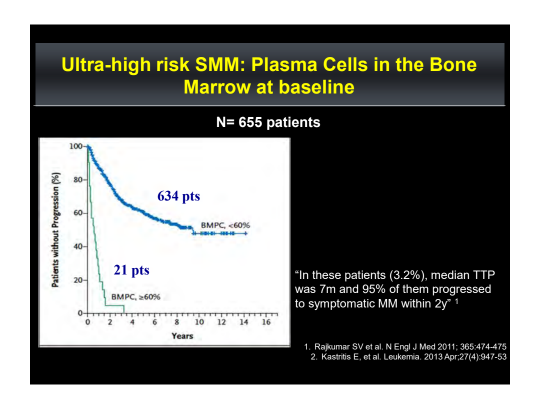
Rajkumar et al. Lancet Oncology 2014; 15: e538-48

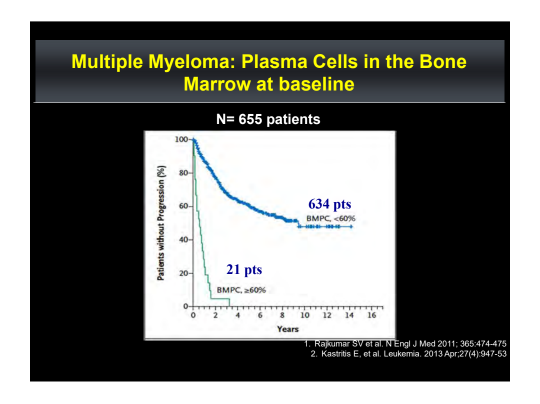
Ultra-high risk SMM: Serum involved/uninvolved free-light chain (FLC) Ratio

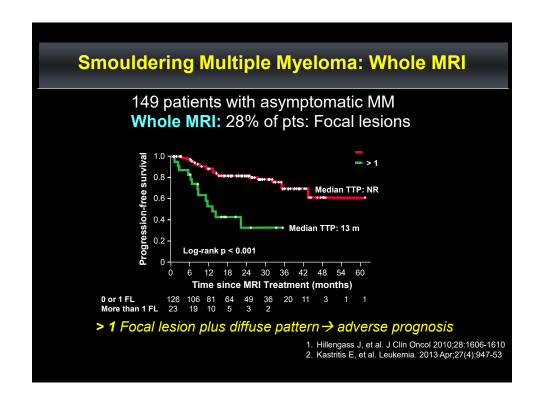


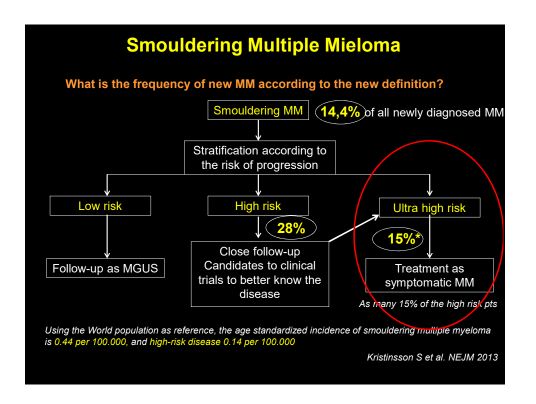
Rajkumar SV et al. N Engl J Med 2011; 365:474-475
 Kastritis E, et al. Leukemia. 2013 Apr;27(4):947-53
 Waxman AJ, et al. J Clin Oncol 32:5s, 2014 (suppl; abstr 8607)



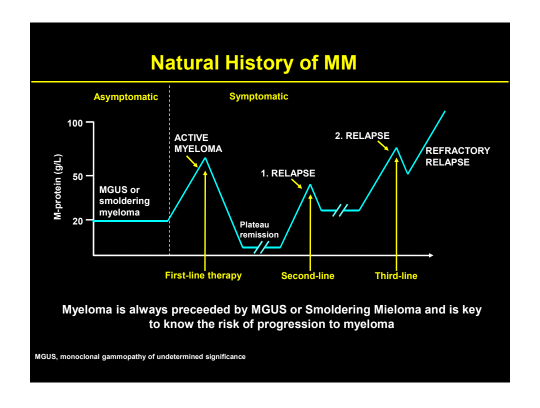


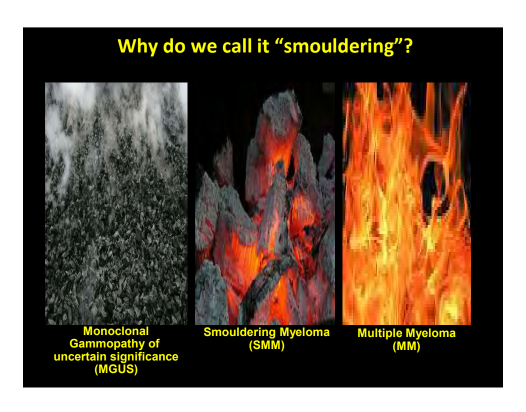






What is the next step? What happens with the other SMM patients?





Transition from MGUS/SMM to MM

- Expansion of altered clones already present in MGUS patients
 López Corral et al. Leukemia 2012
- Branching model→ Key molecular events leading to disease evolution→ distinct patterns of driver mutations

Walker et al. Nature Reviews Cancer 2012

· Differences in inmune surveillance

Dosani et al. Blood Cancer J. 2015

MGUS, monoclonal gammopathy of undetermined significance

Objectives

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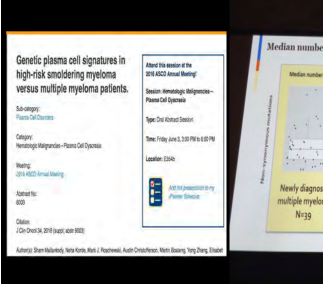
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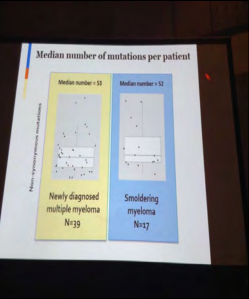
Dosani et al. Blood Cancer J. 2015

We do not know the key mechanism of transition

MGUS, monoclonal gammopathy of undetermined significance

Transition from MGUS/SMM to MM





Transition from MGUS/SMM to MM

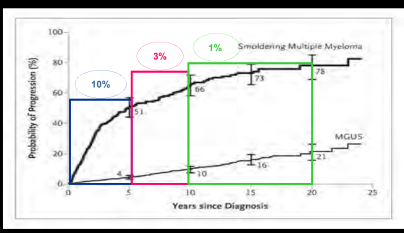




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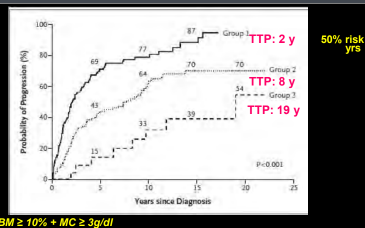




According to the heterogeneity in the risk of progression to MM, we have to identify the individual risk for each new SMM patient.

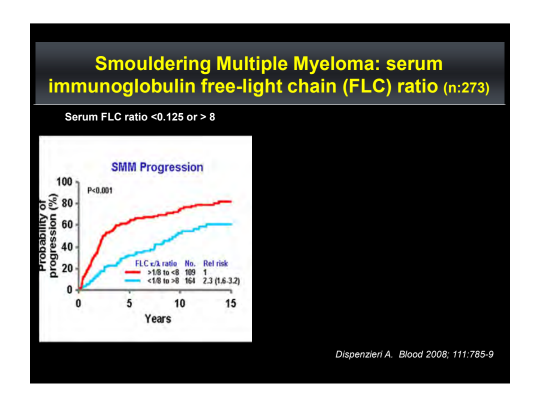
Kyle R. N Engl J Med 2007; 356:2582-90

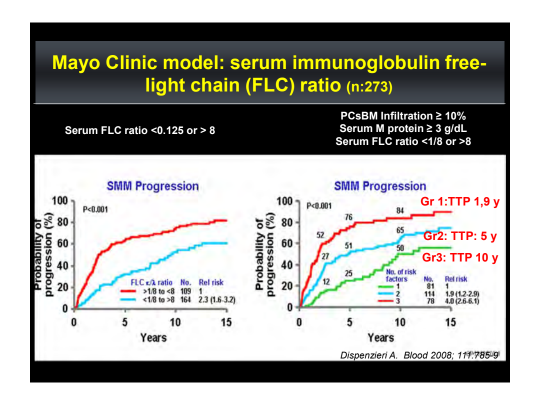
Mayo risk model: PCs BM infiltration and **Serum M-component level**

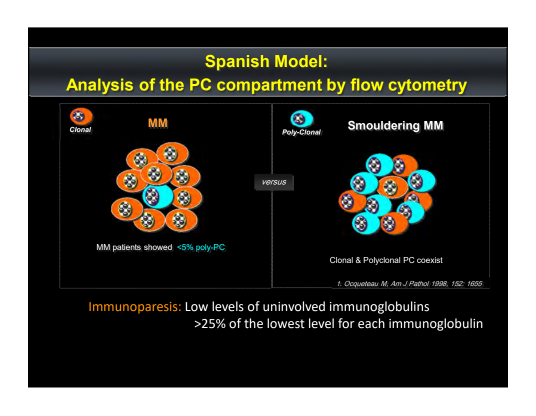


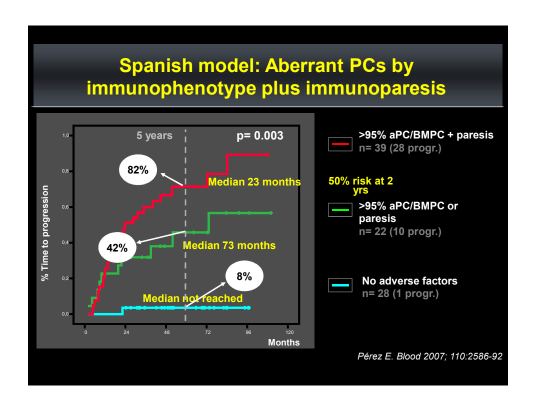
Group 1: PCBM ≥ 10% + MC ≥ 3g Group 2: PCBM ≥ 10% + MC < 3g Group 3: PCBM < 10% + MC ≥ 3g

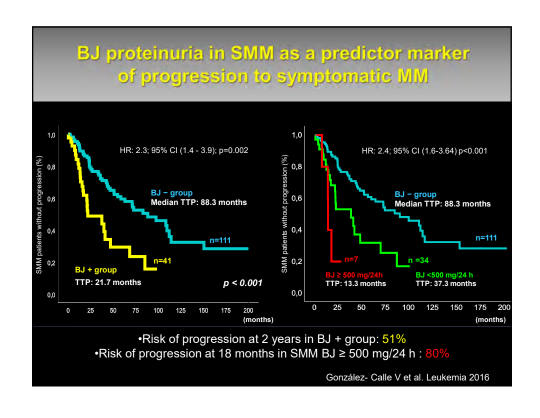
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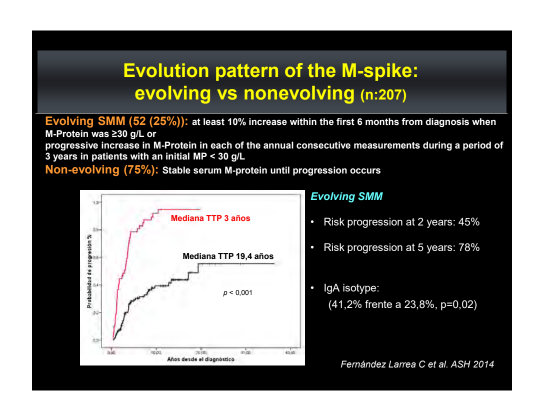






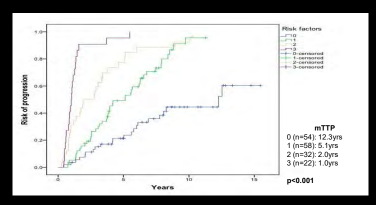






Evolution pattern of the M-spike + eHb + BMPC: (n:190)

Risk factors predicting high risk: 1) eMP as Larrea et al.; 2) eHb: decrease of ≥ 0.5g/dL Hb within 12m of diagnosis; and 3) BMPC infiltratation : ≥20%



SMM with eMP and eHb (with or without BMPC ≥20% had >80% risk of progression to MM within 2 years of diagnosis→ ultra high risk SMM

Ravi et al. ASCO 2016

Del(17p), t(4;14), and +1q21 predict progression from smouldering to symptomatic MM (n=248)

• del(17p13), t(4;14), +1q21 showed significant impact on TTP

	TTP	P
All pts	4.9 years	
+1q21 versus no gain of 1q21	3.7 years 5.3 years	0.013
del(17p13) versus no del(17p13)	2.7 versus 4.9 years	0.019
t(4;14) versus no t(4;14)	2.9 versus 5.2 years	0.021
HD versus NHD	3.9 versus 5.7 years	0.036

- Multivariate analysis: t(4;14), +1q21, HD, reduction of uninvolved immunoglobulins and risk score defined by Kyle et al. as independent factors for adverse outcome
- Conclusion: specific chromosomal aberrations drive transition from asymptomatic to symptomatic disease

Neben et al. JCO 2013; October 21 Epub ahead of print

Primary molecular cytogenetic abnormalities and risk of progression in SMM (n=351)

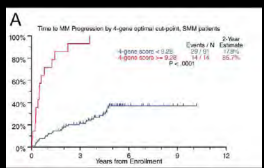
• del(17p13), t(4;14), trisomies showed significant impact on TTP

Cytogenetic abnormalities	TTP
High-risk subgroup	
t(4;14), del(17p)	24 months
Intermediate-risk subgroup	
Trisomy (ies) withouth IgH translocation	34 months
Standard/low-risk subgroup	
T(11;14), other, or no abnormalities	55 months/NR

Rajkumar SV. Leukemia 2013; 27(8): 1738-44

Gene Expression Profiling of purified CD138+ tumor cells in SMM (n: 105)

The validated 70-gene model (GEP-70) identified SMM patients with GEP70>-0.26 with a 51% of progression risk at 2 yrs.

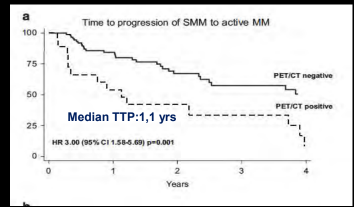


A gene signature derived from 4 genes at an optimal binary cut-point of 9.28, identified 14 patients (13%) with a 2-year therapy risk of 85.7%

Dhodapkar MV et al. Blood 2013 Khan RC et al. Haematologica 2015

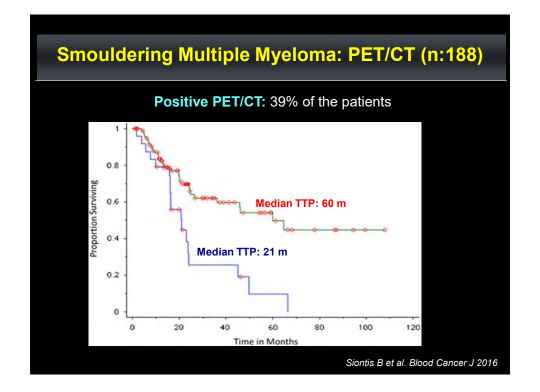
PET-CT in SMM patients as predictor of progression to symptomatic MM (n: 120)

16% of patients had PET positive: 56% of them had 1 FL with a median PET SUV of 4.45 and no osteolysis was observed.



Relative risk of skeletal progression was 3.0 (95% CI 1.3-12, P= 0.013)

Zamagni E et al. Leukemia 2016



Smouldering Multiple Myeloma: Risk models

Identification of high risk SMM→ 50% of progression risk at 2y

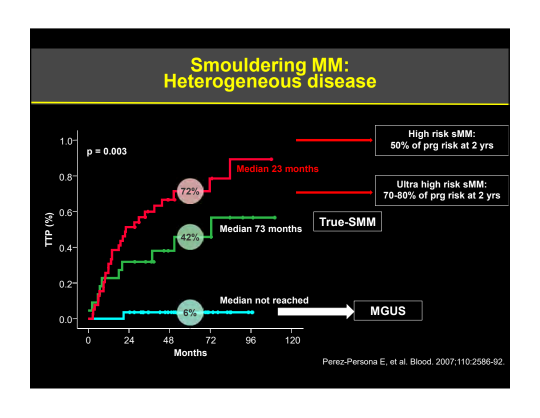
- Mayo Clinic: ≥10% clonal plasma cell bone marrow infiltration, and ≥30g/L of serum M-protein, and serum-free light ratio >0.125 or <8
- Spanish: ≥95% of aberrrant plasma cells measured by flow plus >25% decrease in one or both uninvolved immunoglogulins
- Heidelberg: Tumor mass defined by Mayo risk model plus t(4;14)/del17p/gains of 1q/
- Japanese: Beta 2-microglobulin ≥ 2.5 mg/L plus M-protein increment rate > 1 mg/dL/day
- SWOG: serum M-protein ≥2 g/dL plus involved free light chain >25 and GEP >-0.26 (71% of risk progression at 2 yrs)
- PENN: ≥ 40% clonal PCBM infiltration plus sFLC ratio ≥ 50 plus Albumin □ 3.5 mg/dL (81% of risk at 2 yrs)
- Czech & Heidelberg: immunoparesis plus serum M-protein ≥ 2.3 g/dL plus involved/uninvolved sFLC > 30 (81% of risk at 2 vrs)
- Barcelona: evolving pattern plus serum M-protein ≥ 3 g/dL plus immunoparesis (80% of risk at 2 yrs)

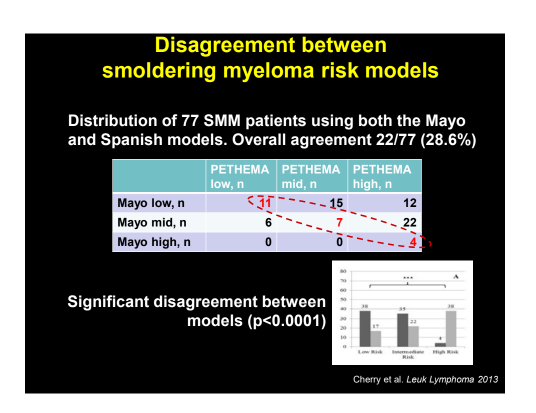
Each model appears to identify patients at high risk, with some but not complete overlap

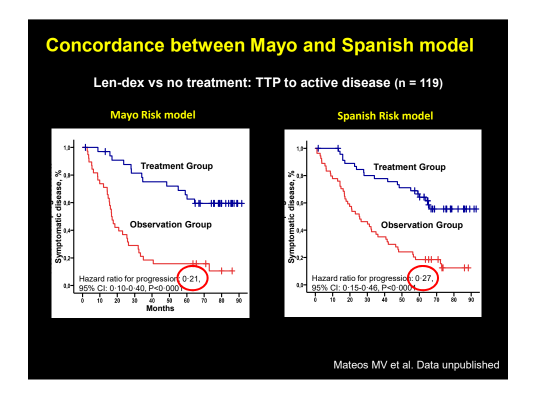
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What is the next step?
What happens for the other SMM patients?

How to proceed once patients are stratified according to the risk?

Smouldering Multiple Myeloma: Management

- Management should be risk-adapted
- Low risk SMM should be followed as MGUS-like pts: annually
- Intermediate risk SMM should be followed as true SMM pts: every 6 months
- Ultra high-risk should be considered MM and be treated
- High-risk SMM can benefit from early treatment

Low and Intermediate risk Smouldering Multiple Myeloma

What is the optimal work-up to do?

- Medical History and physical examination
- Hemogram
- · Creatinine and calcium values
- Protein studies
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- 24-h urine protein electrophoresis (urine M-protein)
- Serum and urine immunofixation
- Serum free light chain mesurement (FLC ratio)
- Bone Marrow aspirate+/- biopsy Only if suspect of active disease
- Skeletal survey/Low-dose CT/PET-CT PET-CT annually for intermediate risk?
- MRI of the spine and pelvis/ Whole-body MRI Repeat at 6 months if one focal lession was present.

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- High-risk SMM can benefit from early treatment

Treatment goals for high-risk smouldering myeloma High-Risk Smoldering Myeloma Progression Early treatment Eradication of Progressive disease Long-term control Cure Advanced disease Chronic disease state Enduring remission © 2011 American Association for Cancer Rese CCR Focus AIR Landgren et al, Clin Cancer Research 2011

Smouldering Multiple Myeloma: Management

Agents	ORR (%)	TTP	os	Reference
Early MP* vs Deferred MP	52 55	No benefit	No benefit	Hjorth M, et al. Eur J Haematol. 1993 Grignani G, et al. Br J Cancer. 1996 Riccardi A, et al. Br J Cancer. 2000
Thal+Zol vs Zol**	37 0	No benefit	No benefit	Witzig TE, et al. Leukemia 2013
Bisphosphonates***vs observation	0	No benefit	No benefit	Martin A, et al. Br J Haematol. 2002 D'arena et al. Leuk Lymphoma. 2011 Musto P, et al. Cancer. 2008

^{*}Abandon: No differences in survival and potential risk of secondary leukemias

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Low, intermediate and high risk patients were included

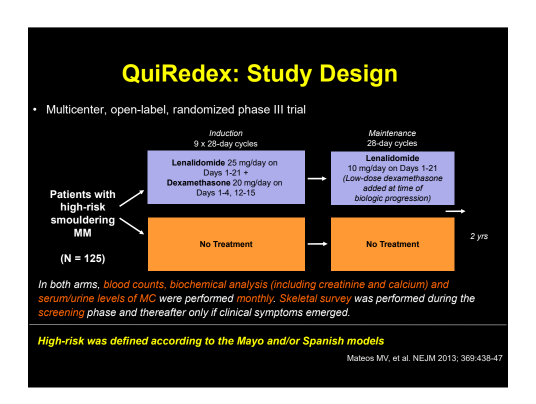
^{**}Low efficacy&high rates of discontinuation due to PN

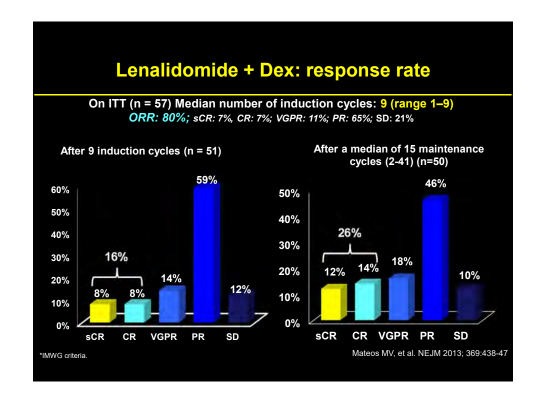
^{***}Skeletal related events lower in the bisphosphonate groups (39% vs 73% and 55% vs 78%)

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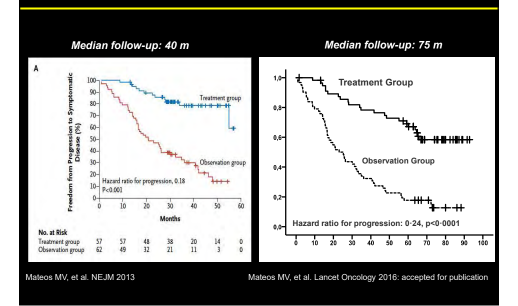
^{**}Low efficacy&high rates of discontinuation due to PN

^{***}Skeletal related events lower in the bisphosphonate groups (39% vs 73% and 55% vs 78%)

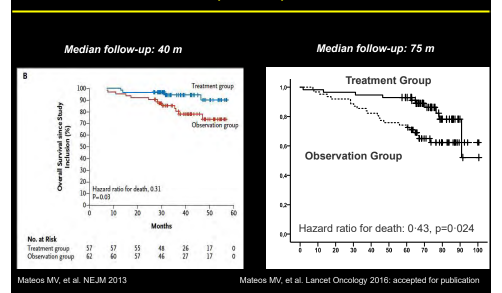




Len-dex vs no treatment: TTP to active disease (n = 119) Per-protocol Patients population



Len-dex vs no treatment: OS from inclusion (n = 119)



Len-dex: biological progressions (n:57 pts)

15 biological progressions during maintenance therapy



Dex was added according to the protocol

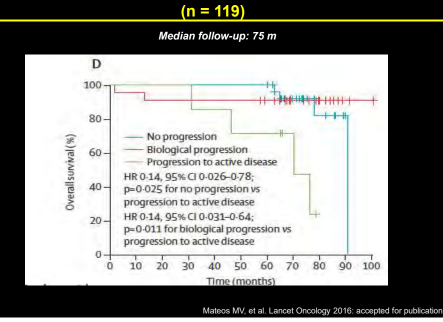
(20 mg 4 days)

- 3 pts achieved PR and 12 stabilized their disease
 - 5 finally progressed to Myeloma
 - 10 pts remain in stable disease and five of them with len 10 mg plus 20 mg dex for 4 days

Why this pre-emptive strategy only for the experimental arm? What was the benefit of this pre-emptive strategy?

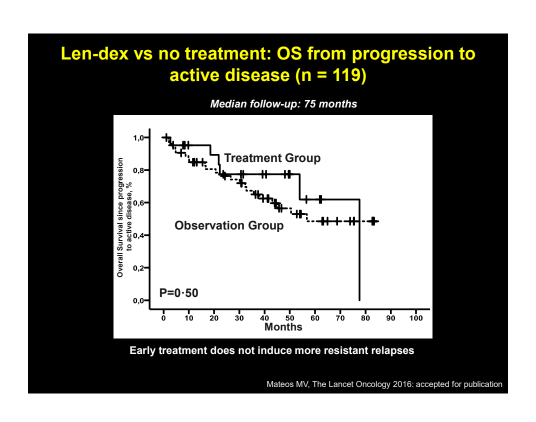
Mateos MV, et al. Lancet Oncology 2016: accepted for publication

Len-dex vs no treatment: OS according to the type of progression



What about rescue therapies	
	9
windi about rescue therables	

Subsequent therapy	Lenalidomide-dex (n=22)	Observation (n=53)
PI-based combinations	13 (59%)	23 (43%)
IMiD's-based combinations	3 (14%)	8 (15%)
PI-IMiD's-based combinations	2 (9%)	16(30%)
Conventional chemotherapy	4 (18%)	6 (11%)
	4 pts (18%) received ASCT	15 pts (28%) received ASCT
	Mateos MV, et al. Lancet Onc	ology 2016: accepted for public



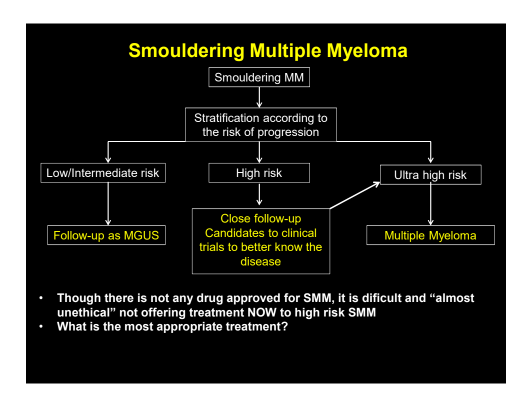
QuiRedex: toxicity profile during induction (n:125)

	Len-dex a	Abstention arm (n:63)	
	G1	G2	G1-2
Anemia	11 (20%)	4 (7%)	2 (4%)
Neutropenia	3 (6%)	8 (14%)	
Thrombopenia	6 (11%)	1 (2%)	
Asthenia	6 (11%)	5 (9%)	6 (11%)
Constipation	4 (7%)	6 (11%)	1 (2%)
Diarrhea	9 (17%)	4 (7%)	2 (4%)
Rash	12 (23%)	6 (11%)	
Infection*	19 (35%)	6 (11%)	14 (26%)
DVT**	1 (2%) 2 (4%)		
SPM -Hematologic -Non hematolog	1 patient (PV) 5 patients*		1 patient (MDS)

^{*3} prostate cancers, 1 breast cancer and 1 cervical epidermoid carcinoma

The cumulative risk of developing a second primary malignancy at 7 years was 12% (95% CI 0á31–11) in the treatment group and 3% (0–4) in the observation group (p=0.070).

Mateos MV, et al. Lancet Oncology 2016: accepted for publication



High-risk Smouldering Multiple Myeloma

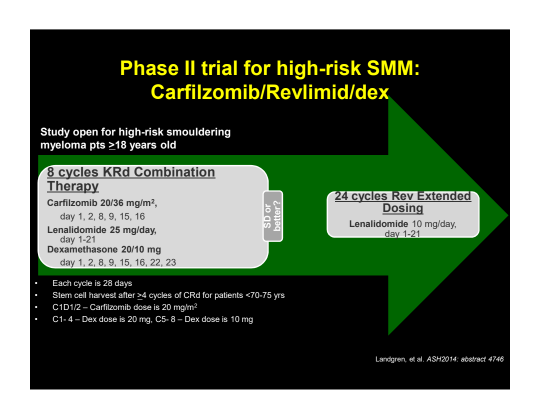
- Len-dex is effective as early treatment, with benefit in TTP to active disease and also in OS
- Numerous clinical trials with several drugs are currently ongoing in this group of patients:

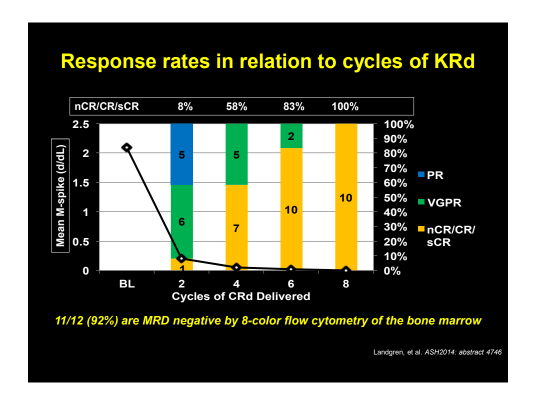
Elotuzumab, daratumumab, Elo-Rd, KRd,

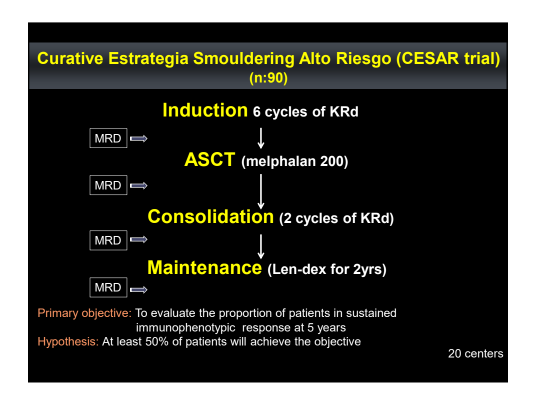
Current Studies in High-Risk Smouldering MM

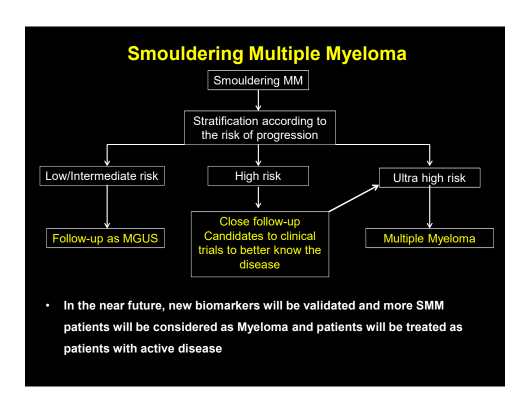
- Biomarker study of elotuzumab (phase II)[2]
- Siltuximab (anti IL6) or no treatment (phase II)[3]
- Biomarker study of BHQ880 (anti DKK1) (phase II)[4]: Data presented at ASH2012: no antitumor effect but anabolic activity
- Lenalidomide or observation (phase III)^[1]
- Elotuzumab-Lenalidomide-dex
- Daratumumab single agent at different doses (Centaurus trial)
- Carfilzomib, lenalidomide, and dexamethasone (phase II)^[5]:

- ClinicalTrials.gov. NCT01169337.
 ClinicalTrials.gov. NCT01441973.
 ClinicalTrials.gov. NCT01484275.
 - ClinicalTrials.gov. NCT01302886.
 ClinicalTrials.gov. NCT01572480.







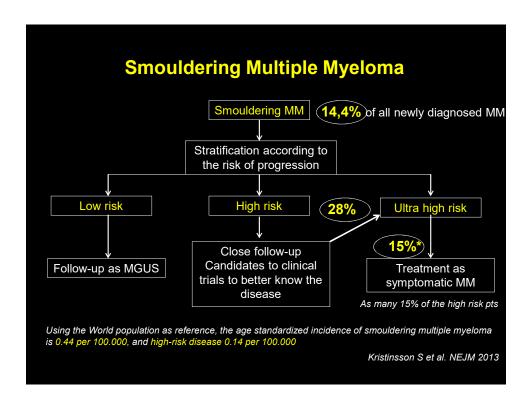


Smouldering Multiple Myeloma: Summary

- The standard of care remains observation until progression to active disease
- The results of the studies would support to plan early treatment in patients at high risk of progression to MM
- Len-dex is effective as early treatment, with benefit in TTP to active disease and also in OS
- Numerous clinical trials with several drugs are currently ongoing in this group of patients

These results support to change the current treatment paradigm for this patient population

Early treatment for high risk SMM patients





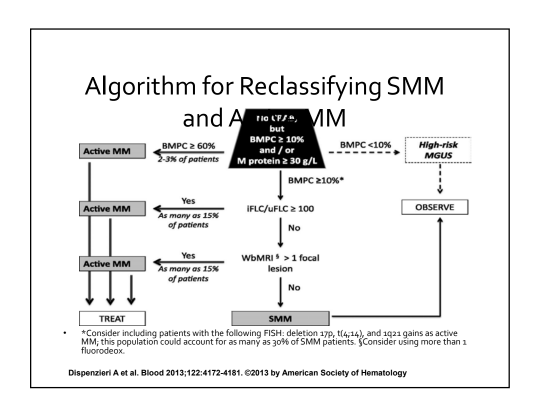
How I treat newly diagnosed transplant eligible patients with multiple myeloma

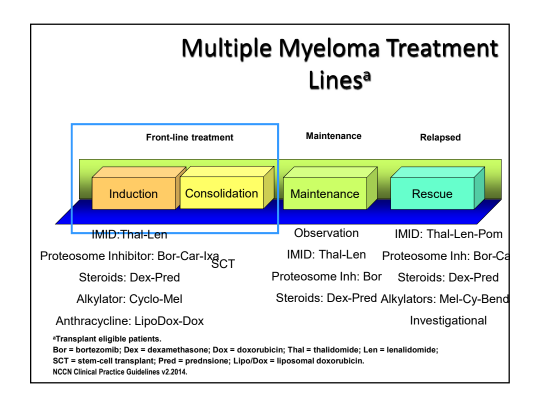
Sergio Giralt MD Melvin Berlin Family Chair in Myeloma Research Professor of Medicine, Weill Cornell Medical College Chief Attending, Adult BMT Service Memorial Sloan Kettering Cancer Center

Start with the basics

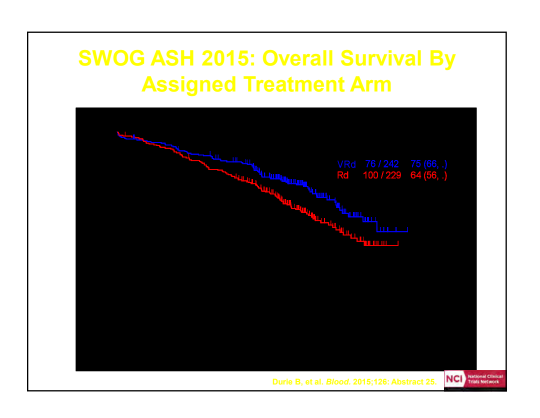
Case Presentation: MJ

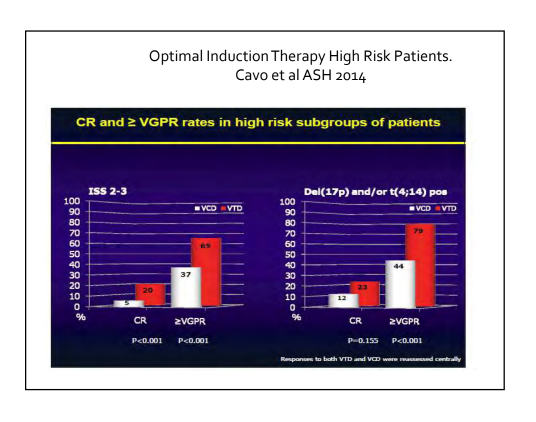
- MJ is a 50 year old female who on routine PE was found with a total elevated serum protein 10.5 gm/lt and a hemoglobin of 11 gm/dl.
- Further work up reveals
- SPEP shows IgG 4,700 mg/dL and kappa 5,200 mg/dL
- M spike 4.2 g/dL
- 24-hour urine was normal < 0.16 g/24 hours
- β₂-microglobulin normal 1.6 mg/L
- Bone marrow biopsy showed 60% plasma cells; normal cytogenetics, no lgH translocations. DECREASED IRON STAINS
- Bone survey showed mild osteopenia

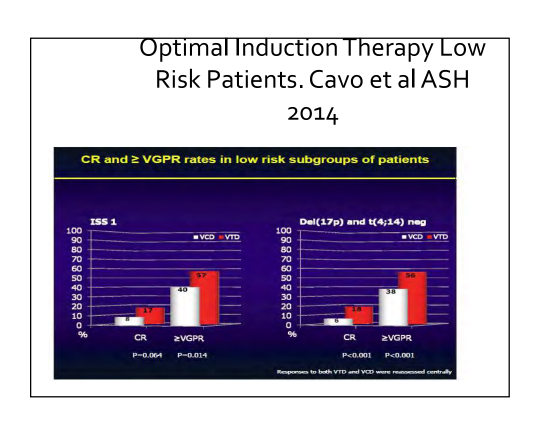


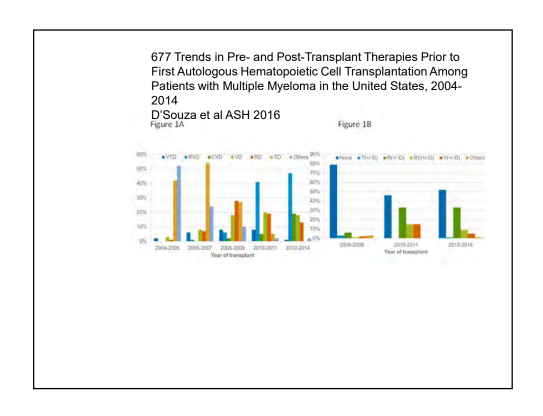


	2015 Confiri Versus Rd (D	
	RVd	Rd
CR	15.7%	8.4%
VGPR	27.8%	23.4%
PR	38%	39.7%
ORR (PR or better)	81.5%	71.5%
SD	15.7%	24.3%
SD or better	97.2%	95.8%
PD or Death	2.8%	4.2%
sessable patients	onal Clinical Network Durie B, et al. Bi	lood. 2015;126: Abstract

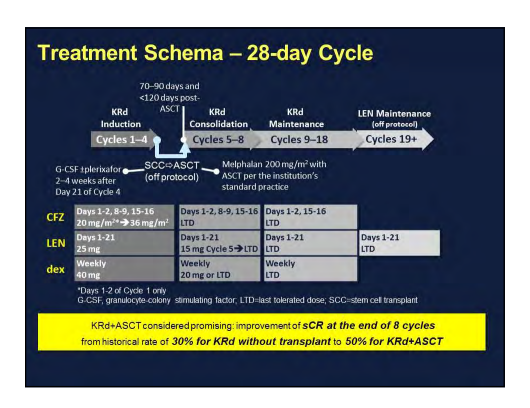


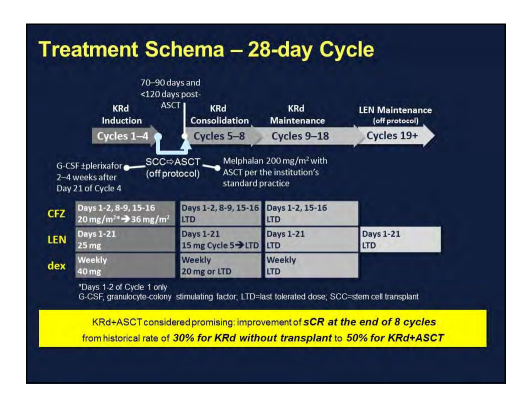






Newer regimens KRD KRD – Dara IXA RD





Daratumumab (DARA) in Combination with Carfilzomib, Lenalidomide, and Dexamethasone (KRd) in Patients (pts) With Newly Diagnosed Multiple Myeloma (MMY1001): an Open-label, Phase 1b Study

Andrzej Jakubowiak,¹ Ajai Chari,² Sagar Lonial,³ Brendan Weiss,⁴ Raymond L. Comenzo,⁵ Kaida Wu,⁵ Nushmia Z. Khokhar,⁵ Jianping Wang,² Parul Doshi,⁶ Saad Z. Usmani⁵

¹University of Chicago Medical Center, Chicago, IL; ²Tisch Cancer Institute, Mount Sinai School of Medicine, New York, NY, USA; ³Department of Hematology and Medical Oncology, Winship Cancer Institute, Emory University, Atlanta, GA, USA; ⁴Abramson Cancer Center and Pereiman School of Medicine, University of Pennsylvania, Philadelphia, PA, USA; ⁵Disnion of Hematology/Oncology, John C. Davis Whyelom and Anyloid Program, Tufs Medical Center, Boston, MA; Pennsylvania, Philadelphia, PA, USA; ⁵Disnion of Hematology/Oncology, John C. Davis Whyelom and Anyloid Program, Tufs Medical Center, Boston, MA; ⁶Janssen Research & Development, LLC, Raritan, NJ, USA; ⁸Levine Cancer Institute/Carcal Roystem, Charlotte, NC, USA; ⁸Disnion State Carcal Roystem, Charlotte

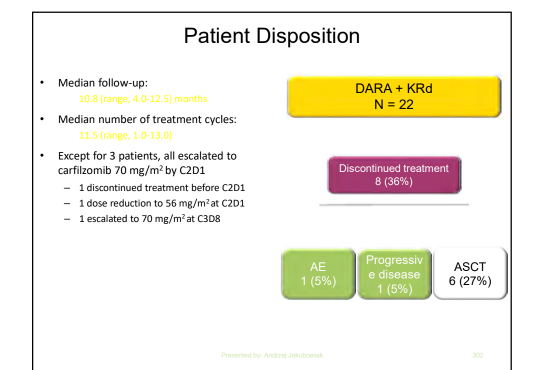
Study Design Dosing Schedule (28-d cycles) **Endpoints** Eligibility/Treatment Daratumumab: Primary Split dose: 8 mg/kg Days 1-2 of Cycle 1 16 mg/kg QW on Cycles 1-2, Q2W on Cycles 3-6, and Q4W thereafter Safety, tolerability •Transplant eligible and noneligible Secondary •Treatment duration: ≤13 Carfilzomib: cycles or until elective discontinuation for ASCT 20 mg/m² C1D1 response, time to response, IRR Escalated to 70 mg/m² C1D8+; weekly (Days 1, 8, 15) •No clinically significant Lenalidomide: Exploratory 25 mg; Days 1-21 of each cycle required at screening Dexamethasone: 40 mg/weeka

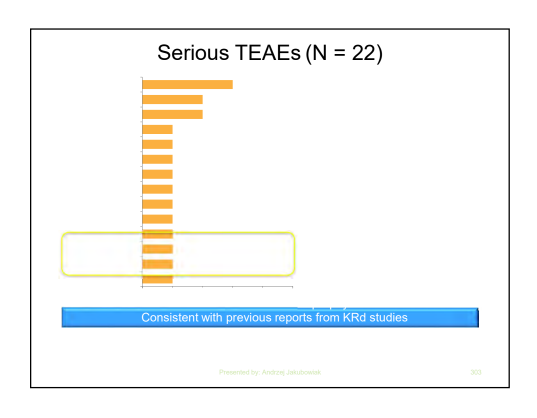
Baseline Demographics

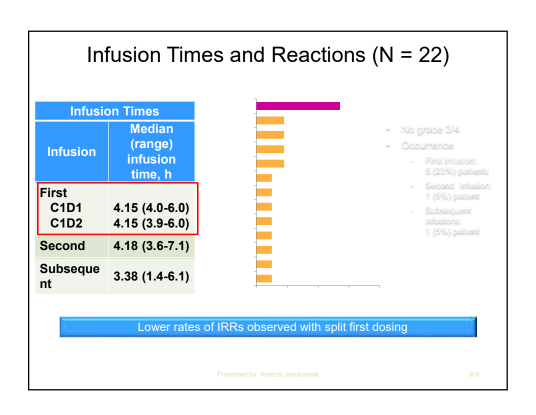
Characteristic	DARA + KRd (N = 22)
Age, years, n (%)	
Median (range)	59.5 (34-74)
<65	15 (68)
65 - <75	7 (32)
Gender, n (%)	
Male	12 (55)
Female	10 (46)
Race, n (%)	
White	19 (86)
African American	1 (5)
American Indian or Alaska Native	1 (5)
Not reported	1 (5)
ECOG score, n (%)	
0	12 (55)
1	9 (41)
2	1 (5)

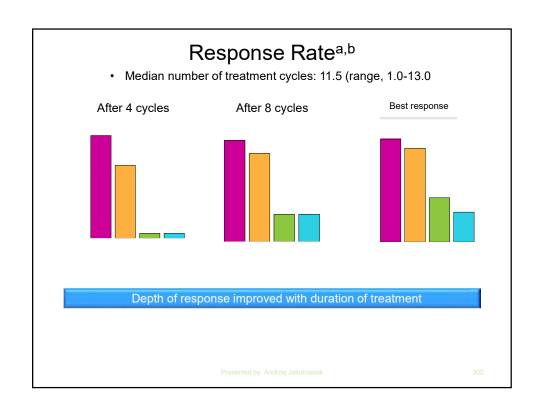
Presented by: Andrzei Jakubowia

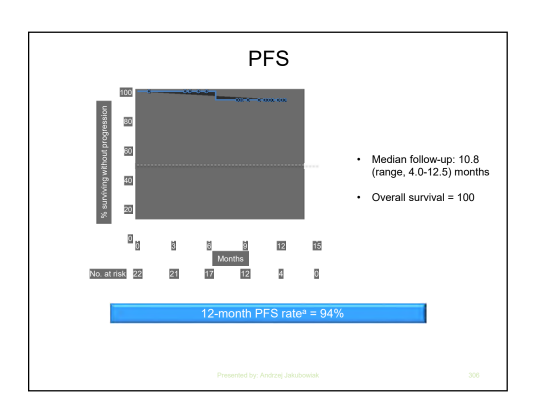
301











Stem Cell Harvest and ASCTa

- Median number of CD34⁺ cells collected from patients: 10.4 x 10⁶ cells/kg (n = 19)
- Median 5 treatment cycles prior to stem cell harvest

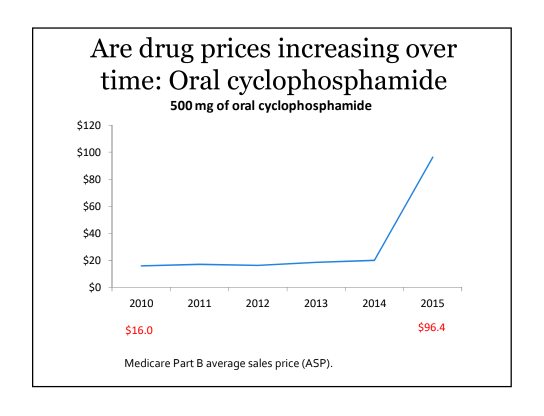
Pat	tient	Stem cell mobilization	Total CD34 ⁺ cells (x10 ⁶ /kg body weight)	Treatment cycle at ASCT	Best response
	1	Plerixafor and Filgrastim	30	9	sCR
:	2	Plerixafor and Filgrastim	12	5	VGPR
:	3	Plerixafor and Filgrastim	28	4	VGPR
	4	Filgrastim	38	4	VGPR
	_	Plerixafor and	40.4	=	KCDD
		Stem cell yield is cons	istent with previou	s KRd stud	ies
	6	Filgrastim	6.5	4	VGPR

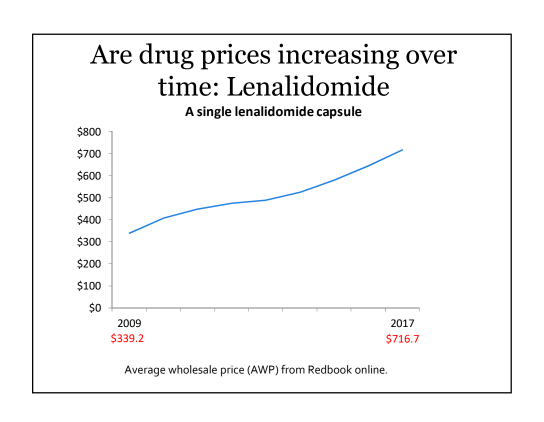
Presented by: Andrzei Jakubowial

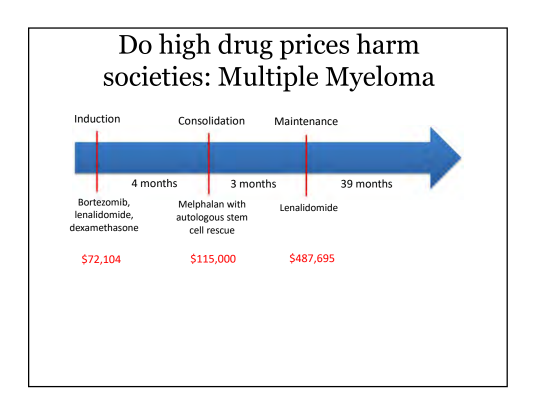
307

Lenalidomide-Ixazomid Dexa (RID) NDMM EHA 2017

	Richardson et al	Kumar et al	Dimopoulos et al
Ixazomib dose	3.0-3.7 Twice Weekly	1.6-3.8 Weekly	4.0 mg weekly CTX 300/400 mg/m2 1-8-15
N	40 (did not proceed to HCT)	42 (did not proceed to HCT)	70 non transplant eligible
ORR	93%	80	
CR+VGPR	68%	63%	
CR	32%	32%	
Median PFS	24.9 m	25 m	56% at 2 years
% Discontinuing due to AE Most Common	NS Rash and Neutropenia	NS Rash and Neutropenia	24% Neutropenia
i			





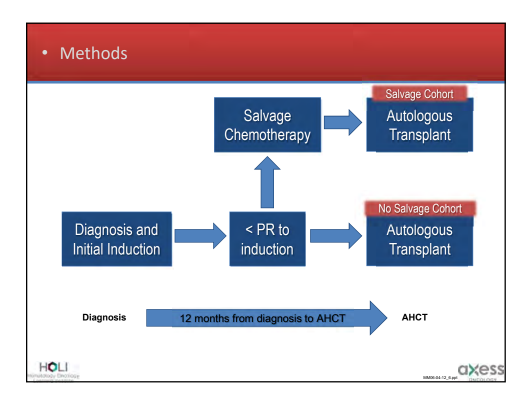


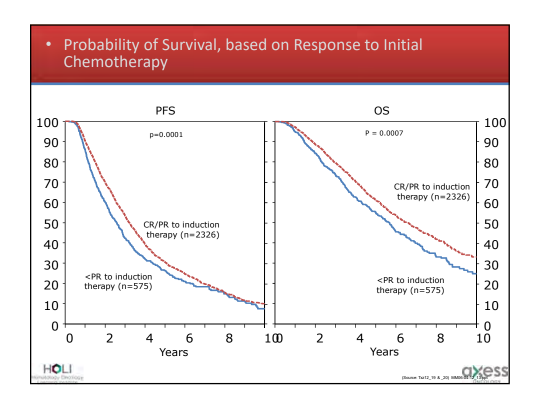
Optimal induction for Latin America

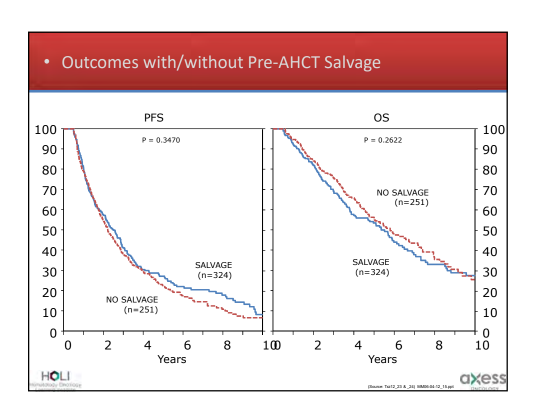
- Most cost effective approach to achieve a major response VGPR or greater
- CY-BOR-D x 4
- If no VGPR after 4 cycles consider
 - RVD x 4

 Effect of Pre-transplant Salvage Therapy Prior to Autologous Transplant (AHCT) in Patients Not Responding to Initial Induction for Multiple Myeloma (MM)

CIBMTR Study MM06-04

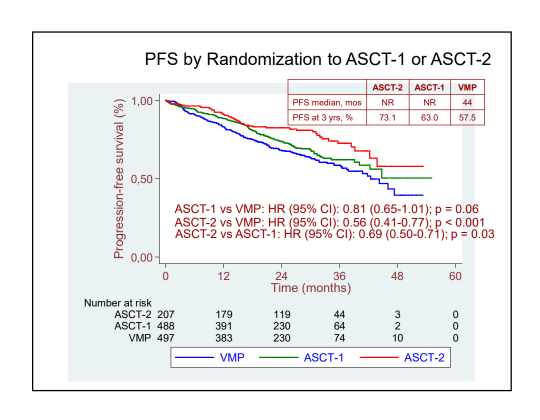


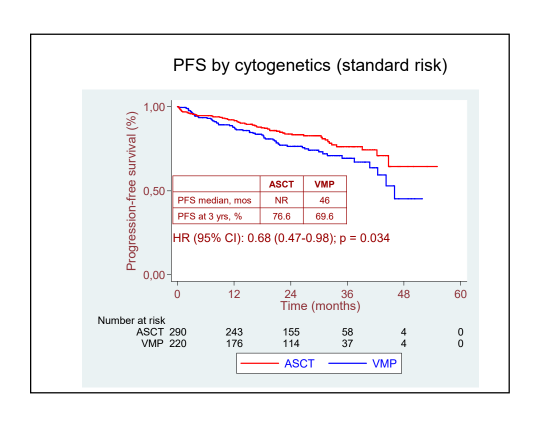


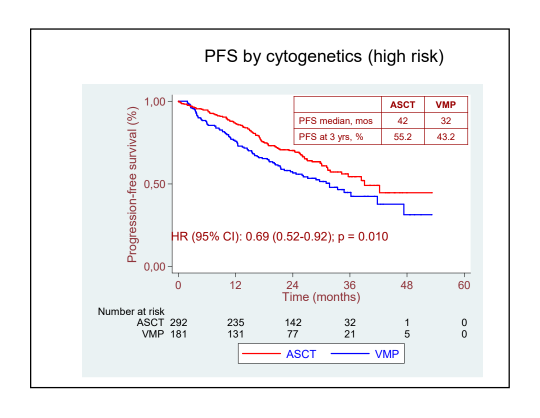


CONSOLIDATION = CONTROVERSY

991 Upfront Single Versus Double Autologous Stem Cell Transplantation for Newly Diagnosed Multiple Myeloma: An Intergroup, Multicenter, Phase III Study of the European Myeloma Network (EMN02/HO95 MM Trial) Cavo et al.





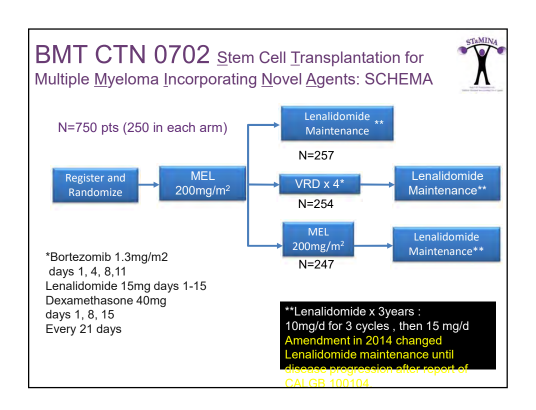


Primary Results from the Randomized Prospective Phase III Trial of the Blood and Marrow Transplant Clinical Trials Network

> (BMT CTN 0702 – STaMINA Trial) NCT#01109004

Autologous Hematopoietic Cell Transplant (AHCT), with and without Consolidation (with Bortezomib, Lenalidomide (Len) and Dexamethasone) and Len Maintenance versus Tandem AHCT and Len Maintenance for Up-Front Treatment of Patients with Multiple Myeloma





BMT CTN 0702: Regimens prior to Transplant

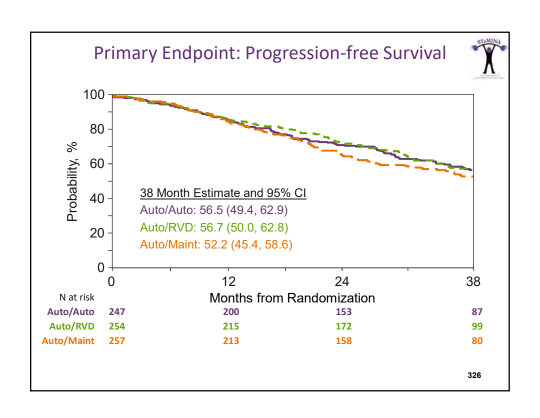


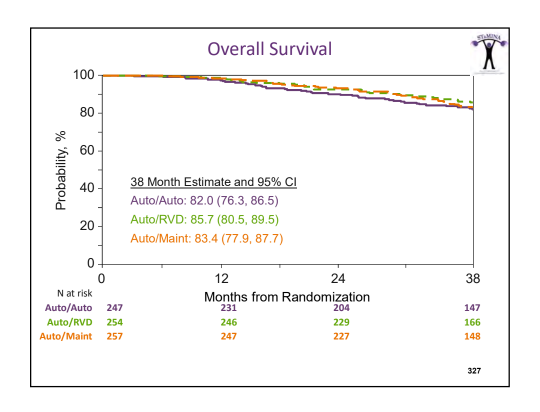
		Auto/Auto (N=247)		Auto/RVD (N=254)		Maint 257)
	N	N %		%	N	%
Initial Therapy						
Bort/Len/Dex	141	57.1	134	52.8	143	55.6
Cy/Bort/Dex	33	13.4	35	13.8	40	15.6
Len/Dex	24	9.7	28	11.0	22	8.6
Bort/Dex	28	11.3	32	12.6	32	12.5
Other	21	8.5	25	9.8	20	7.8

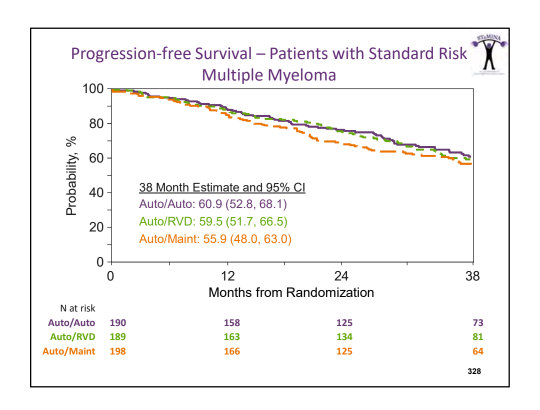
Bort, bortezomib; Cy, cyclophosphamide; Dex, dexamethasone; Len, lenalidomide

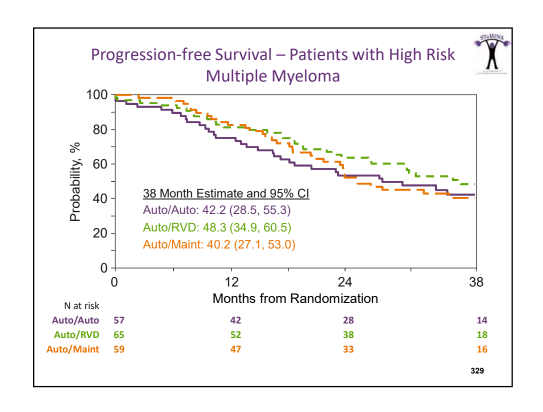
324

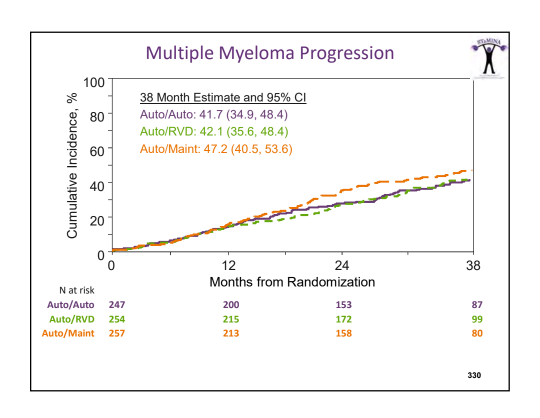
Compliance with each interventic						
	Auto// (N=2			/RVD 254)	1	Maint 257)
	N	%	N	%	N	%
Received ^{2nd} Intervention						
No	79	32.0	30	11.8	-	-
Yes	168	68.0	224	88.2	-	-
Started maintenance						
No	41	16.6	43	16.9	14	5.4
Yes	206	83.4	211	83.1	243	94.6
						325

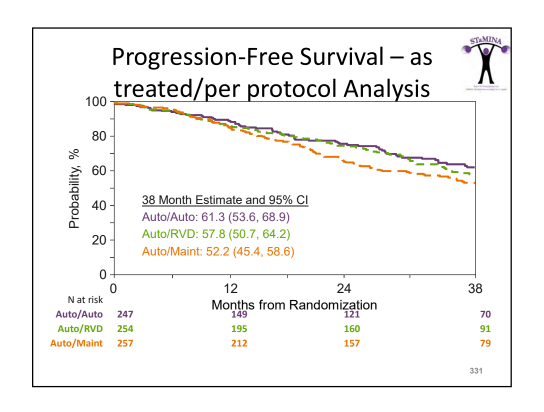












Conclusion



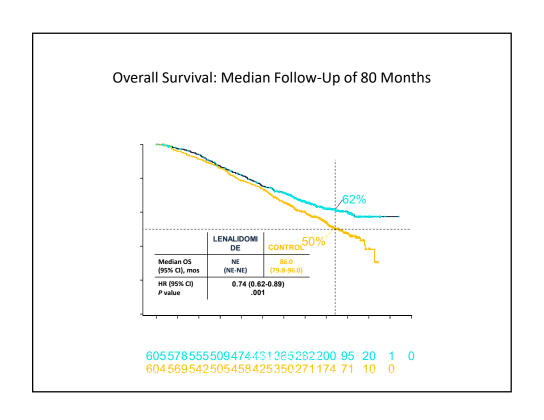
 In the era of thalidomide analogues and proteasome inhibitors used in the initial therapy for myeloma (in this study >90% either, >50% both) and the use of prolonged maintenance therapy with lenalidomide, post transplant consolidation with cycles of RVD or a second transplant do not produce incremental PFS benefit.

332

Lenalidomide Maintenance After High-Dose Melphalan and Autologous Stem Cell Transplant in Multiple Myeloma: A MetaAnalysis of Overall Survival

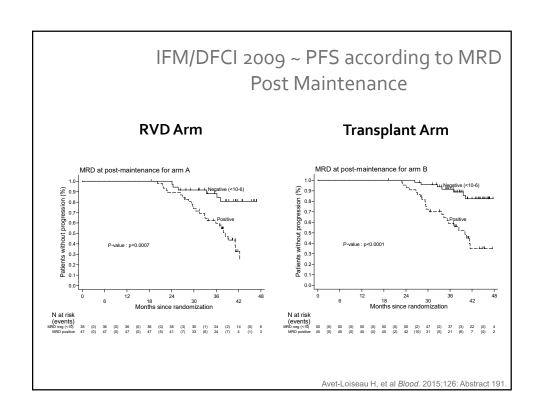
Michel Attal,¹ Antonio Palumbo,² Sarah A. Holstein,³
Valérie Lauwers-Cances,¹ Maria Teresa Petrucci,⁴ Paul Richardson,⁵ Cyrille
Hulin,⁶ Patrizia Tosi,⁶ Kenneth C. Anderson,⁵ Denis Caillot,⁶ Valeria
Magarotto,⁶

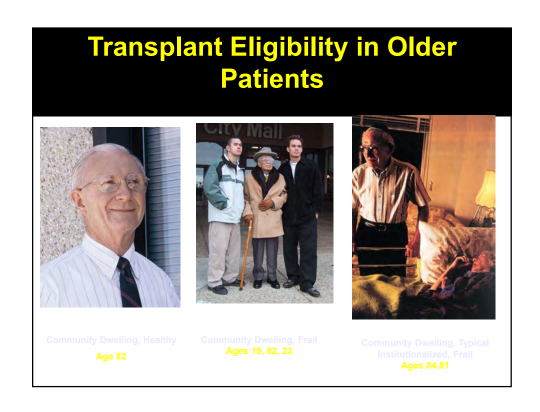
Philippe Moreau, ¹⁰ Gerald Marit, ¹¹ Zhinuan Yu, ¹² Philip L. McCarthy ¹³

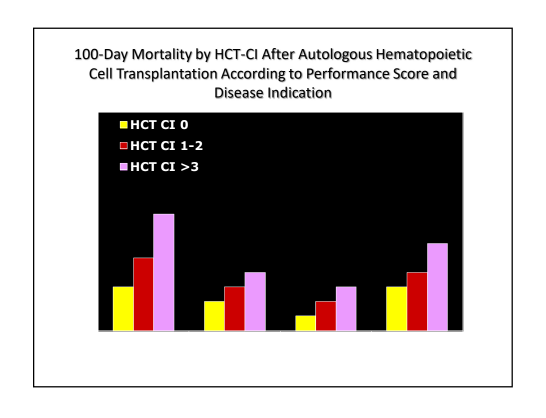


All SPMs After Randomization

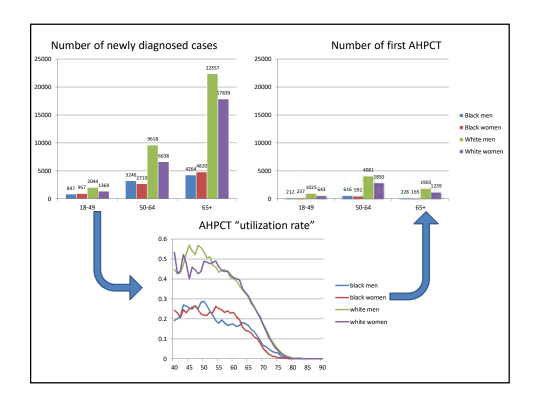
CALGBa		LGB ^a	IF	М	GIMEMA	
SPMs, n ^b	LEN (n = 224)	Placeb o (n = 221)	LEN (n = 306)	O (n = 302)	LEN (n = 56)	No MNTC (n = 79) ^c
Hematologic AML ^d MDS B-cell malignancy Other	15 7 4 4 0	8 0 4 3 1	21 6 4 11 1	9 3 3 2 1	0 - - - -	0 - - - -
Solid tumors	17	10	21	13	5	2







CLINICAL TRIALS AND OBSERVATIONS Geriatric assessment predicts survival and toxicities in elderly myeloma patients: an International Myeloma Working Group report Table 2. The final Cox regression model HR (95% CI) Score Antonio Palumbo, ¹ Sara Bringhen, ¹ Maria-Victoria Matess, ² Alessandra Larocca, ¹ Thierry Facon, ³ Shaji K. Kumar, ⁴ Age, y ≤75 76-80 Massimo Offidari, ⁵ Philip McCarthy, ⁶ Andrea Evangelista, ⁷ Sagar Lonial, ⁸ Sonja Zweegman, ⁹ Pelegrino Musto, ¹⁰ 1.13 (0.76-1.69) .549 Evangelos Terpos, ¹¹ Andrew Belch, ¹² Roman Hajek, ¹³ Heinz Ludwig, ¹⁴ A. Keith Stewart, ¹⁵ Philippe Moreau, ¹⁶ 2.40 (1.56-3.71) <.001 Kenneth Anderson, ¹⁷ Hermann Einsele, ¹⁸ Brian G. M. Durie, ¹⁹ Meletios A. Dimopoulos, ¹¹ Ola Landgren, ²⁰ 1.67 (1.08-2.56) .020 Jesus F. San Miguel, 21 Paul Richardson, 22 Pieter Sonneveld, 25 and S. Vincent Rajkurrar⁴ IADL 1.43 (0.96-2.14) CCI 1.37 (0.92-2.05) .125 1 2.37 (1.38-4.09) 3.21 (1.85-5.58) Favorable Unfavorable 1.79 (1.23-2.60) Missing 1.13 (0.69-1.83) .036 Therapy Proteasome inhibitors 0.74 (0.50-1.11) 142 HRs and relative risks are for OS in patients with the factors as compared with those without the factors. The model was adjusted for ISS, chromosome abnormalities, and therapy. Unfavorable profile defined as 1(4;14) or 1(14;16) or 6417213. abnormalness, and strong, del17p13. AIC = 1748.918; Harrell C index = 0.7069.



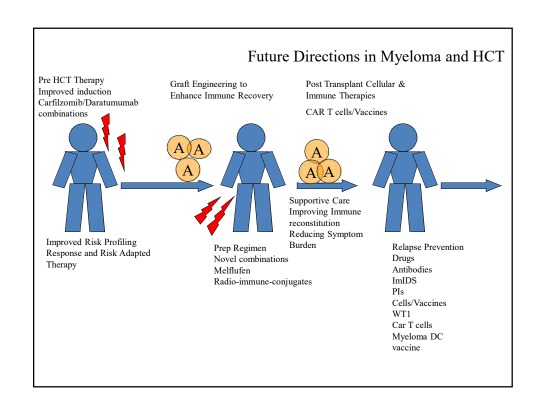
1190 Significant Differences in Stem Cell Transplant Utilization Rates (STUR) of Autologous Hematopoietic Cell Transplant (AHCT) in Multiple Myeloma (MM) Based on Ethnicity without Differences in Efficacy. a CIBMTR Report. D'Souza et al.

Table 1. Stem cell Transplant utilization rate estimate

Year	Hispanic (95%CI)	Non-Hispanic White (95%CI)	Non-Hispanic Black (95%CI)
2008	8.6 (7.9 -9.4)%	22.6 (21.8 -23.9)%	12.2 (11.4 -13.0)%
2009	9.8 (9.0 -10.7)%	26.6 (25.7 -27.5)%	13.2 (12.4 -14)%
2010	11.9(10.9-13.0)%	29.4(28.4-30.4)%	15.7(14.8-16.8)%
2011	11.4(10.6-12.4)%	34 (32.9 - 35.1)%	18.2(17.1-19.3)%
2012	14.2(13.1-15.4)%	35.4(34.3-36.6)%	19(18-20.2)%
2013	16.9(15.6 -18.3)%	37.8 (35.5 - 38)%	20.5(19.4-21.8)%

Table 2. Outcomes of AHCT in MM by ethnicity (values are expressed as probabilities with 95% confidence intervals)

Outcome	H (N=1933)	NHW (N=18046)	NHB (N=4123)	p-value
TRM				0.4
100 days	0.6 (0.3-1)%	0.9 (0.7-1)%	0.6 (0.4-0.9)%	0.2
1 year	2 (2-3)%	3(2-3)%	3 (2-3)%	0.7
PFS				1.0
1-year	82 (80-84)%	83 (82-83)%	82 (81-83)%	0.3
2-year	66 (64-68)%	66 (65-67)%	66 (64-67)%	0.9
3-year	54 (51-56)%	53 (52-54)%	54 (52-55)%	0.8
OS				0.1
1-year	94 (93-95)%	94 (93-94)%	94 (94-95)%	0.3
2-year	86 (85-88)%	86 (85-86)%	86 (85-87)%	0.7
3-year	80 (77-82)%	77 (77-78)%	79 (77-80)%	0.05



Frontline Treatment for MM Transplant Eligible

- Current standard is
 - Triple induction (ImID,PI, steroids)
 - Single high dose melphalan auto HCT
 - Maintenance lenalidomide
- HCT severely underutilized in United States
- Risk Stratification is still NOT STANDARD
- However...
 - Patients with high risk features reasonable to explore
 - IMID/PI maintenance
 - Tandem auto or auto/allo HCT
- MRD Directed therapy not yet standard but all roads point towards that destination

Treatment of the Fit/Older MM Patient





Myelomacenter.org run9001@med.cornell.edu

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Disclosures for Ruben Niesvizky

In compliance with ACCME policy, following disclosures to the session audience:

Research Support/P.I.	Celgene, Takeda, Amgen, Janssen, BMS
Employee	N/A
Consultant	Celgene, Takeda, Amgen, Janssen, BMS
Major Stockholder	N/A
Honoraria	N/A
Speakers Bureau/Scientific Advisory Board	N/A

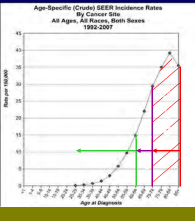
The Elderly Patient

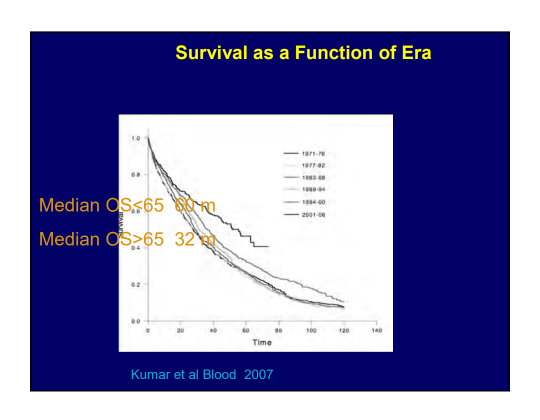
The median age at diagnosis is 70 years

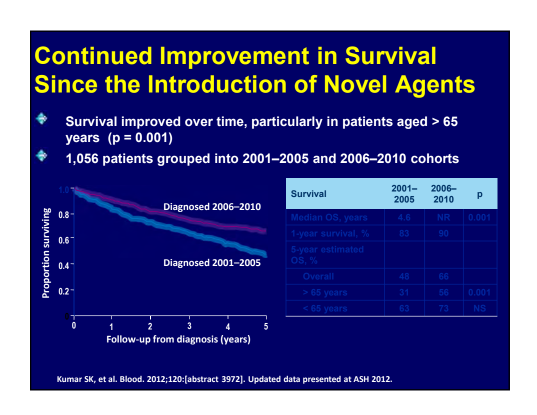
- Nearly half of multiple myeloma patients are considered elderly
- ▼ Traditionally the definition of elderly based on transplant eligibility
 (European and North American trials)
 - Patients under 65 years of age, 35%
 - Older patients from 65 to 75 years

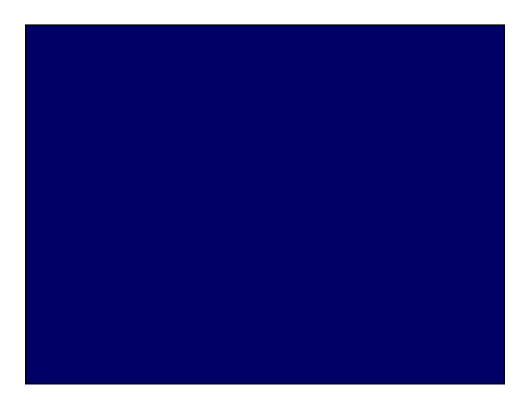
of age, 28%

Palumbo A, et at Herrico Gartier Scottoward Educ Program. 2009:566-577.; Ferlay J, et al. GLOBOCAN 2002 Garcel Incidence, Mortality and Prevalence Worldwide. IARC CancerBase No. 5 Version 2.0. Lyon: IARC Press; 2004.; Ries LAG, et al. National Cancer Institute. SEER Cancer Statistics Review. Source: SEER 13. Accessed August 24, 2010 at: http://seer.cancer.gov/faststats



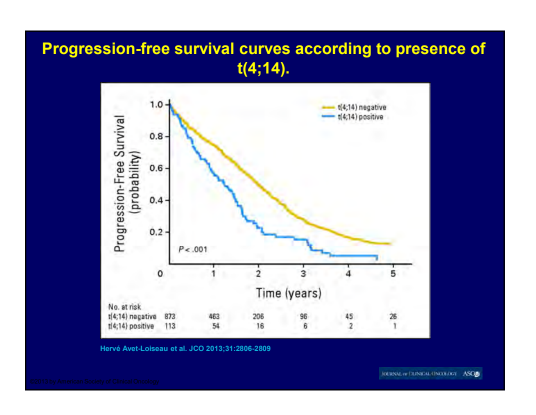


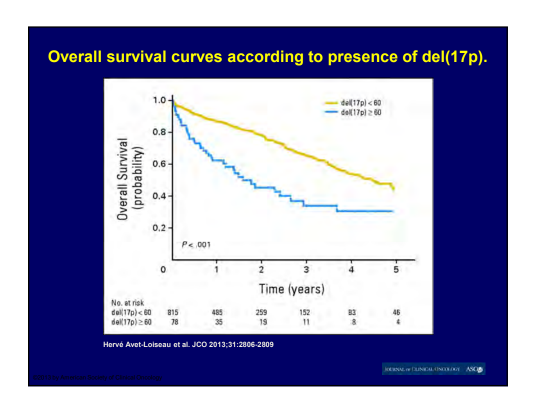




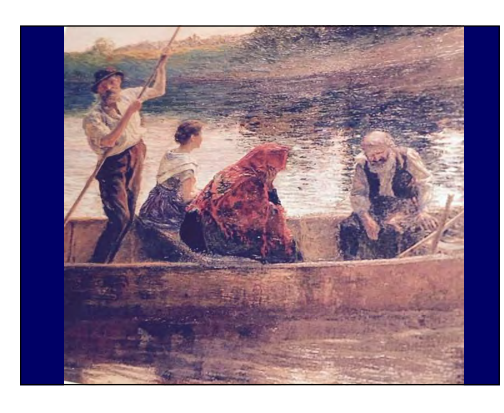
- Biology of the Disease
 - FISH and ploidy
 - GEP
 - Others
- ♦ Host
 - Comorbidities
 - Functional status
- Therapeutics and Toxicology
 - Response and dose modifications
- Psychosocial Aspects
 - Access to care and social support

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2-Year Mortality Rate for Persons Age 70 Years and Older

- * 8% if fully independent
- ♦ 14% if dependent in IADL
- ♦ 27% if dependent in ADL
- >40% if institutionalized

Comorbidity Is a Key Factor in Survival

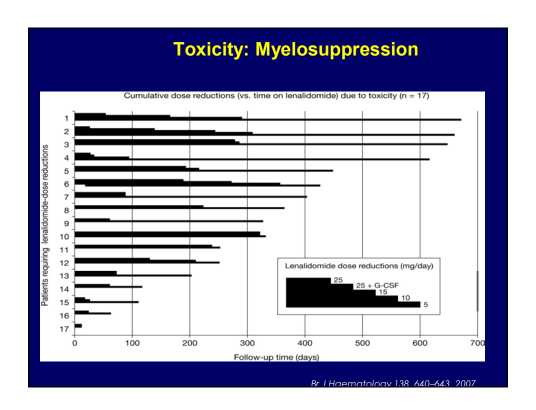
Age-Comorbidity		Actual 10-Year
Score	N	Survival (%)
0-1	369	97-99
2	136	87
3	109	79
4	42	47
5	29	34

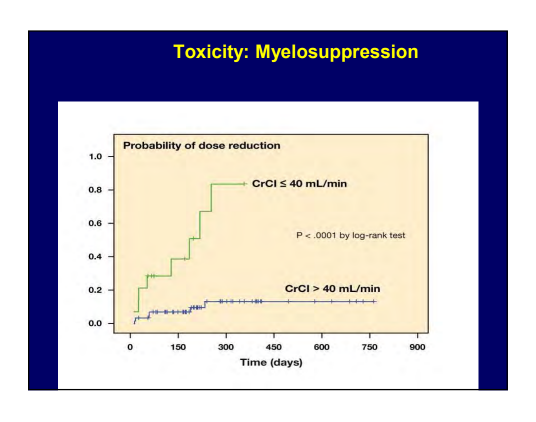
Charlson et al. J Chronic Dis. 1987;40:373.

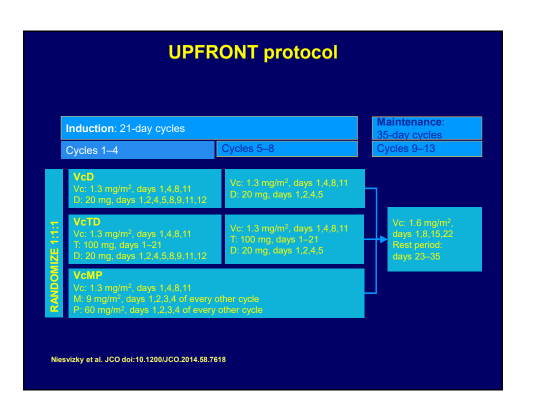
Aspecto	Método	Conclusión Clínica y Proporcionalidad
Funcional	Motilidad, Desempeño físico, Fatiga,	Inversamente or Directamente supervivencia
Psiquiátrico	HADS	Directamente proporcional a la morbilidad psiquiática
Social	MOS 3MS	Inversamente or Directamente supervivencia
Nutrición	NMA	Supervivencia
Co-morbilidad	Formula calculada	Supervivencia

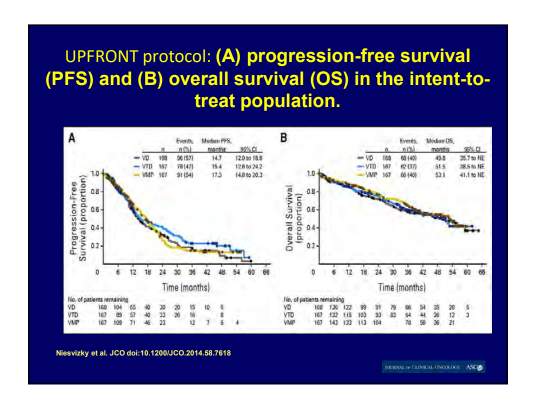
- Biology of the Disease
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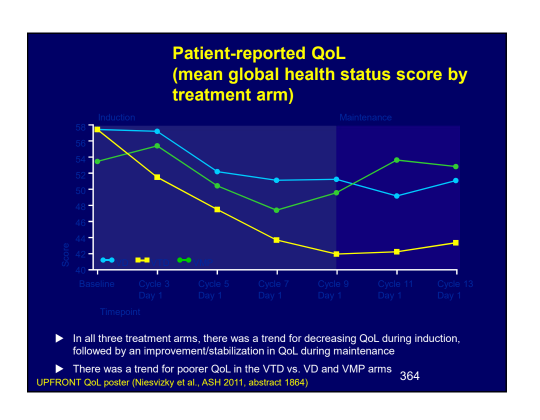
P450 CYP Renal Clearance











Once-weekly administration of bortezomib as a strategy to improve tolerability

Study details	Grade ¾ Gl toxicity	Grade 3/4 peripheral neuropathy	Discontinuation due to AE
VISTA: VMP ¹⁻³ Bortezomib twice-weekly	20%	14%	34%
(GIMEMA) ⁴ Bortezomib once-weekly	-	5%	17%
(PETHEMA/GEM) ⁵ Bortezomib once-weekly	7%	7%	12% [†]

Once-weekly administration of bortezomib as a strategy to maintain/improve the efficacy

Study details	CR+PR	CR	PFS	3 yrs-OS
VISTA: VMP ¹⁻³ Bortezomib twice-weekly	71%	30%	TTP:24 m	68%
Modified VISTA ⁴ (GIMEMA) Bortezomib once-weekly VMPT->VT VMP	90% 81%	42% 24%	37 m 27 m	85% 80%
Modified VISTA ⁵ (PETHEMA) Bortezomib once-weekly VMP vs VTP→VT vs VP	80%	23% → 42%	31 m	70%

^{1.} San Miguel et al. NEJM 2008;359:906 2. San Miguel et al. NEJM 2008;359:906; Supplementary Appendix 3. Mateos et al. J Clin Oncol 2010;28:2259-66

^{4.} Palumbo et al. JCO 2010; 28:5101-095. Mateos et al. Lancet Oncol 2010;11:934-41

^{1.} San Miguel et al. NEJM 2008;359:906 2. San Miguel et al. NEJM 2008;359:906; Supplementary Appendix 3. Mateos et al. J Clin Oncol 2010;28:2259-66

^{4.} Palumbo et al. J Clin Oncol 2010;28:5101-9 5. Mateos et al. Lancet Oncol 2010;11:934-41

Bortezomib IV versus SC

222 relapsed and/or refractory MM patients. Bz is given at conventional dose and scheme

	Bortezomib IV (n=73)	Bortezomib SC (n=145)
Primary +/-dex))	endpoint: response after 4/8	cycles (single agent bortezomib or
ORR	42%/52%	42%/52%
CR	8%/12%	6%/10%
TTP	9∙4 m	10·4 m

	Bortezo	mib IV	Bortezo	mib SC
	All grades	Grade ≥3	All grades	Grade ≥3
Periph Neurop	53%	16%	38%	6%
				P=0·04 and 0·03

No diferences in pharmakokinetics studies

Moreau et al. Lancet Oncology 2011; 12(5): 431-40 Arnulf B et al. Haematologica 2012: Epub ahead of print

- Biology of the Disease
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Regular Article

CLINICAL TRIALS AND OBSERVATIONS

Geriatric assessment predicts survival and toxicities in elderly myeloma patients: an International Myeloma Working Group report

Antonio Palumbo, ¹ Sara Bringhen, ¹ Maria-Victoria Mateos, ² Alessandra Larocca, ¹ Thierry Facon, ³ Shaji K. Kumar, ⁴ Massimo Offidani, ⁵ Philip McCarthy, ⁸ Andrea Evangelista, ⁷ Sagar Lonial, ⁸ Sonja Zweegman, ⁹ Pellegrino Musto, ¹⁰ Evangelos Terpos, ¹¹ Andrew Belch, ¹² Roman Hajek, ¹³ Heinz Ludwig, ¹⁴ A. Keith Stewart, ¹⁵ Philippe Moreau, ¹⁶ Kenneth Anderson, ¹⁷ Hermann Einsele, ¹⁸ Brian G. M. Durie, ¹⁹ Meletios A. Dimopoulos, ¹¹ Ola Landgren, ²⁰ Jesus F. San Miguel, ²¹ Paul Richardson, ²² Pieter Sonneveld, ²³ and S. Vincent Rajkumar⁴

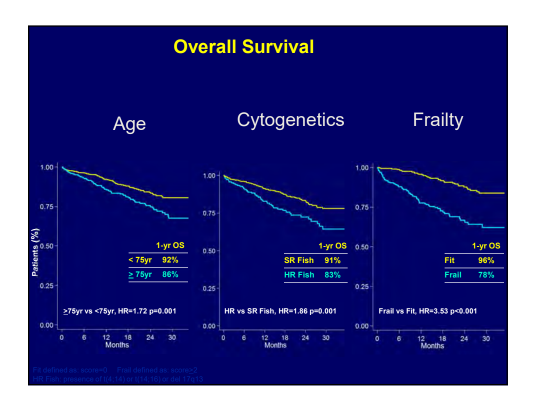


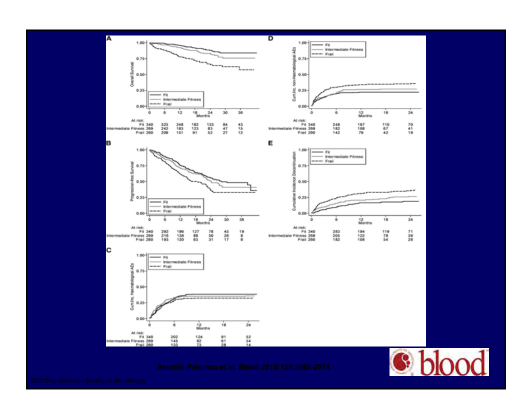
Leading the way in experimental and clinical research in hematology

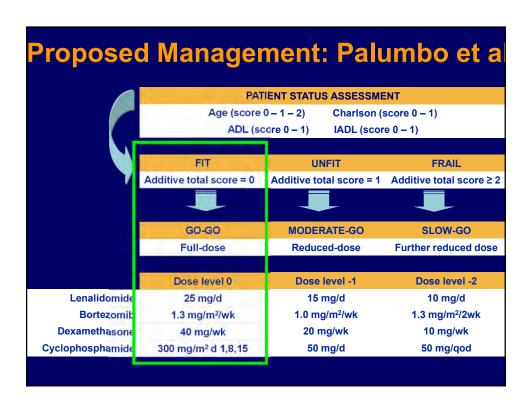
March 26, 2015; Blood: 125 (13)

	Frailty	Index		
Variable		HR (CI 95%)	Р	SCORE
AGE	Age <75 years	1	-	0
	Age 75-80 years	1.37 (0.93-2.03)	0.114	1
	Age >80 years	2.75 (1.81-4.18)	<0.001	2
CHARLSON INDEX	Charlson <u><</u> 1	1	-	0
	Charlson <u>></u> 2	1.6 (1.07-2.39)	0.021	1
ADL SCORE	ADL >4	1	-	0
	ADL <u><</u> 4	1.76 (1.14-2.71)	0.01	1
IADL SCORE	IADL >5	1	_	0

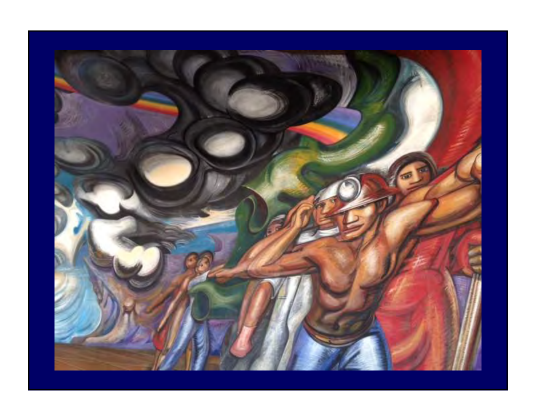
ADDITIVE TOTAL SCORE	PATIENT STATUS
0	FIT
1	UNFIT
<u>≥</u> 2	FRAIL

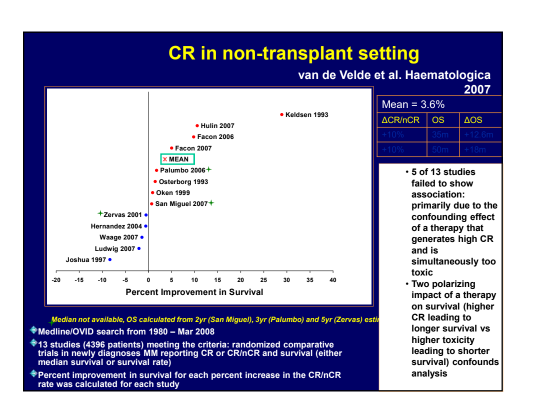






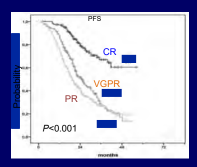


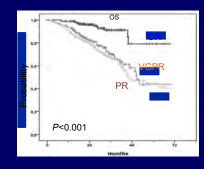




Hematologic CR correlates with long-term PFS and OS in elderly patients treated with novel agents

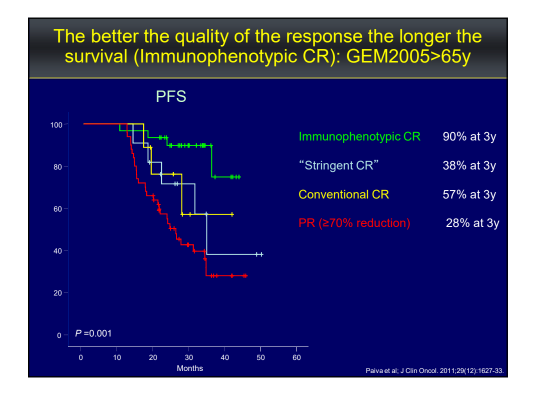
- Retrospective analysis: 3 randomized European trials of GIMEMA and HOVON groups (N=1175)
- First-line treatment
 MP (n=332), MPT (n=332), VMP (n=257), VMPT-VT (n=254)





*Significant benefit also seen when analysis is restricted to patients >75 years old

Gay et al. Blood 2011; 117(11):3025-31

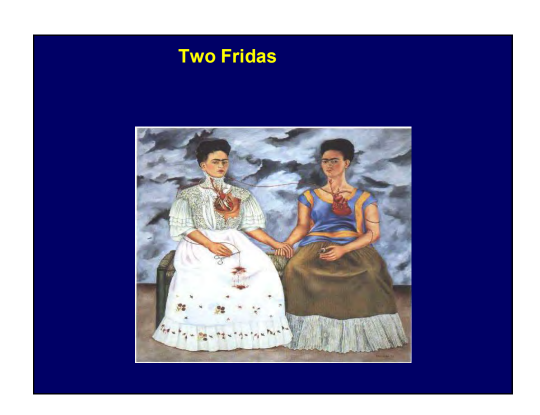


The Older Fit Patient

Therapeutic Considerations:

Is CR/MRD the goal?

Is Transplant the way?



Older Patients: To Transplant or Not

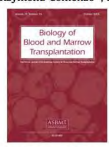
- Performance status and/or functional status
- Cardiac function
- Pulmonary function
- Liver function
- Infectious disease
- Psychosocial support
- Patient goals and preferences

AuPBSC Transplants Age ≥60 yrs at time of HCT P trend YEARS 94-95 04-05 96-97 98-99 00-01 02-03 39 127 179 529 955 1337 OS d100 94 92 94 97 98 97 0.45 (82 -(90-90-97) (95-(96-(97-99)98) 98) 98) 98) OS 12 73 79 91 (90- <.00 84 92 92 month (57-(71-(78-(90-(90-93) 1 post tx 86) 85) 90) 93) 93)

Cost-Effectiveness of Autologous Hematopoietic Stem Cell Transplantation for Elderly Patients with Multiple Myeloma using the Surveillance, Epidemiology, and End Results-Medicare Database



Gunjan L. Shah ^{1,*}, Aaron N. Winn ^{2,3}, Pei-Jung Lin ², Andreas Klein ⁴, Kellie A. Sprague ⁴, Hedy P. Smith ⁴, Rachel Buchsbaum ⁴, Joshua T. Cohen ², Kenneth B. Miller ⁴, Raymond Comenzo 4, Susan K. Parsons



Biology of Blood and Marrow Transplantation

Volume 21, Issue 10

	Transplantation	Nontransplantation
Living more than 2 years	n = 234	n = 180
First year after diagnosis	\$8337	\$2607
Middle years	\$2435	\$2088
Last year	\$8114	\$6809
Living less than 2 years	n = 36	n = 90
Monthly	\$13,106	\$6756

Total cost of care per month during each time frame. Significant differences were seen only in the first year after diagnosis for patients living longer than 2 years and monthly for those living less than 2 years.

Age Is Not a Prognostic Variable With Autotransplants for Multiple Myeloma

By D.S. Siegel, K.R. Desikan, J. Mehta, S. Singhal, A. Fassas, N. Munshi, E. Anaissie, S. Naucke, D. Ayers, D. Spoon, D. Vesole, G. Tricot, and B. Barlogie

Multiple myeloma (MM) typically afflicts elderly patients with a median age of 65 years. However, while recently shown to provide superior outcome to standard treatment, high-dose therapy (HDT) has usually been limited to patients up to 65 years. Among 550 patients with MM and a minimum follow-up of 18 months, 49 aged ≥65 years were identified (median age, 67; range, 65 to 76 years). Their outcome was compared with 49 younger pair mates (median, 52; range, 37 to 64 years) selected among the remaining 501 younger patients (<65 years) matched for five previously recognized critical prognostic factors (cytogenetics, β₂-microgic C-reactive protein, albumin, creatinine). Nearly one half had been treated for more than 1 year with standard therapy and about one third had refractory MM. All patients received high-dose melphalan-based therapy; 76% of the younger and high-cose melphalan-based interapy, reach this year-gas and 65% of the older group completed a second transplant (P=.3). Sufficient peripheral blood stem cells to support two HDT cycles (CD34 $> 5 \times 10^6 / \mathrm{kg}$) were available in 83% of younger and 73% of older patients (P=.2). After HDT, hematopoletic recovery to critical levels of granulocytes

(>500/µL) and of platelets (>50,000/µL) proceeded at comparable rates among younger and older subjects with both first and second HDT. The frequency of extramedullary toxicities was comparable. Treatment-related mortality with the first HDT cycle was 2% in younger and 8% among older subjects, whereas no mortality was encountered with the second transplant procedure. Comparing younger/older subjects, median durations of event-free and overall survival were 2.8/1.5 years (P = .2) and 4.8/3.3 years (P = .4). Multivariate analysis showed pretransplant cytogenetics and B2microglobulin levels as critical prognostic features for both event-free and overall survival, whereas age was insignificant for both endpoints (P=.2/.8). Thus, age is not a biologically adverse parameter for patients with MM receiving high-dose melphalan-based therapy with peripheral blood stem cell support and, hence, should not constitute an exclusion criterion for participation in what appears to be superior therapy for symptomatic MM.

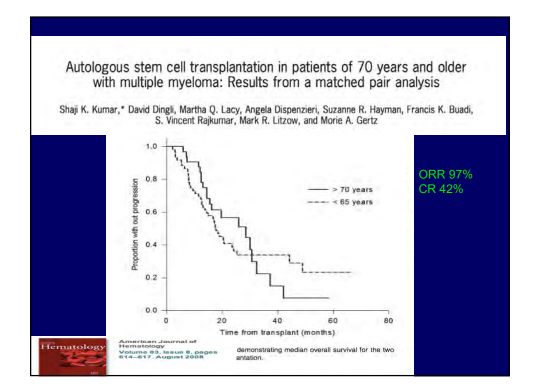
1999 by The American Society of Hematology.

Autologous stem cell transplantation in elderly multiple myeloma patients over the age of 70 years

ASHRAF BADROS, BART BARLOGIE, ERIC SIEGEL, CHRISTOPHER MORRIS, RAMAN DESIKAN, MAURIZIO ZANGARI, ATHANASIOS FASSAS, ELIAS ANAISSIE, NIKHIL MUNSHI AND GUIDO TRICOT Myeloma and Transplantation Research er University of Arkansas for Medical Sciences Little Rock AR HSA

P < .001.

P = .013.



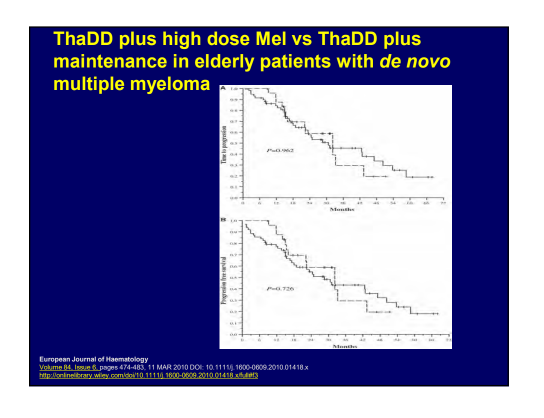
ORIGINAL ARTICLE

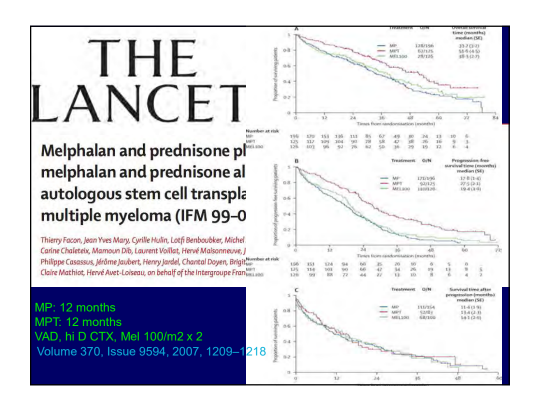
ThaDD plus high dose therapy and autologous stem cell transplantation does not appear superior to ThaDD plus maintenance in elderly patients with *de novo* multiple myeloma

Massimo Offidani¹, Pietro Leoni¹, Laura Corvatta², Claudia Polloni¹, Silvia Gentili¹, Agnese Savini¹, Francesco Alesiani², Marino Brunori², Massimo Catarini², Giuseppe Visani², Arduino Samori², Maurizio Burattini², Riccardo Centurioni², Mauro Montanari¹, Paolo Fraticelli², Miriana Ruggieri², Sadia Falcioni², Piero Galieni²

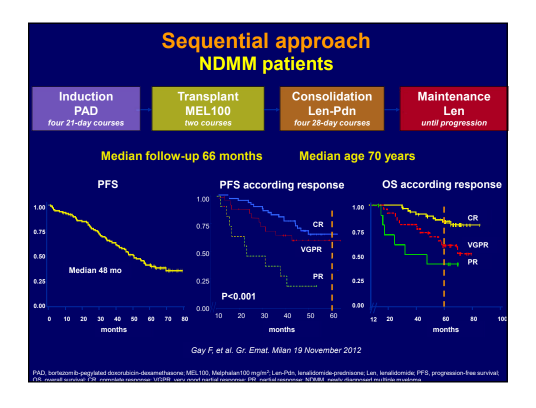
¹Clinica di Ematologia Azienda Ospedaliero-Universitaria, Ospedali Riuniti di Ancona, Ancona; ²Marche Myeloma Network (GEMaMM), Ancona, Italy







Francesca Sara Pezza Paola Ome	Gay, ¹ Valeria tti, ⁵ Samanth dé, ¹ Vittorio N	Magarotto, ¹ Cla a Perrari, ² Anna Montefusco, ⁹ Ma	ion-maintenance audia Crippa, ² Norbert Pesc a Marina Liberati, ⁶ Stelania aria Teresa Petrucci, ¹⁰ Nicol oro, ¹ Paolo Corradini, ⁹ and	costa, ³ Tommasina Guglie Oliva, ¹ Francesca Patrian la Giuliani, ¹¹ Roberto Pas	Imelli, ⁴ Federica Cavallo, ¹
Induc PA four 21-da	ction D	Tra M	insplant C	Consolidation Len-Pdn our 28-day courses	Maintenance Len until progression
	PAD	PAD+ MEL	PAD+MEL+Len	+ Len maint	
CR	12%	78%	48%	53%	
VGPR	43%	43%	32%	29%	
PR	33%	17%	14%	13%	
SD	11%	6%	5%	4%	
PD	0	0	© bloo	Leading the	



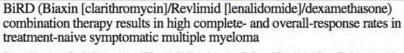
The Older Fit Patient

Therapeutic Considerations:

Is CR/MRD the goal?

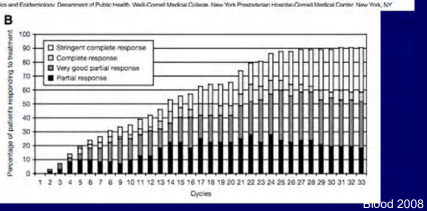
Is transplant the way?

Is continued treatment best?

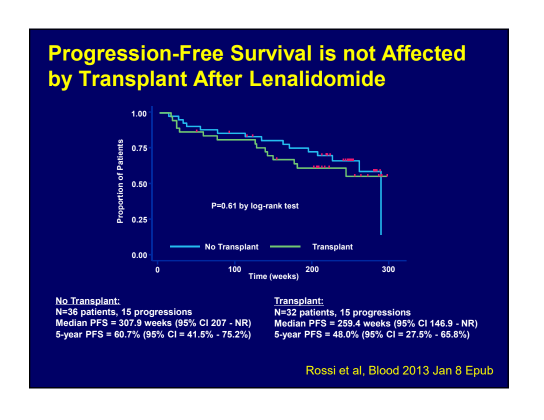


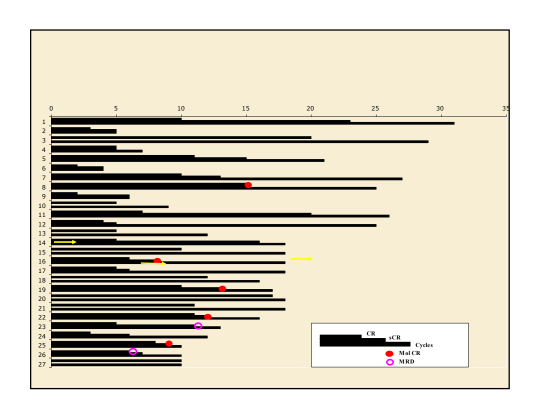
Ruben Niesvizky,¹ David S. Jayabalan,¹² Paul J. Christos,³ Jessica R. Furst,¹ Tara Naib,¹ Scott Ely,² Jessica Jalbrzikowski,¹ Roger N. Pearse,¹ Faiza Zafar,¹ Karen Pekle,¹ April LaRow,¹ Richard Lent,² Tomer Mark,¹ Hearn J. Cho,¹ Tsiporah Shore,¹ Jeffrey Tepler,¹ John Harpel,¹ Michael W. Schuster,¹ Susan Mathew,² John P. Leonard,¹ Madhu Mazumdar,³ Selina Chen-Kiang,² and Morton Coleman¹

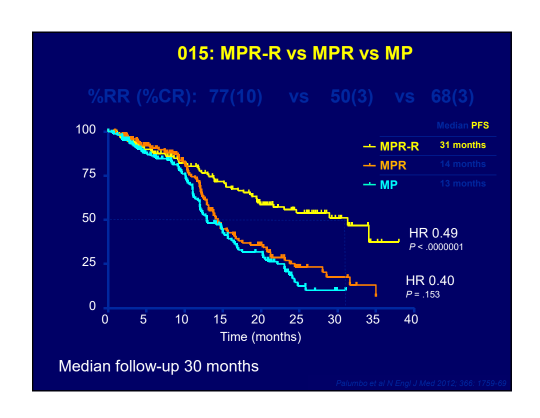
*Center of Excellence for Lymphoma and Myeloma, Division of Hematology and Medical Oncology, Department of Medicine, *Department of Pathology, and *Division of Biostatistics and Ecidemiology. Department of Public Health. Well-Cornell Medical College. New York Presbytarian Hoscibal-Cornell Medical Context. New York, NY

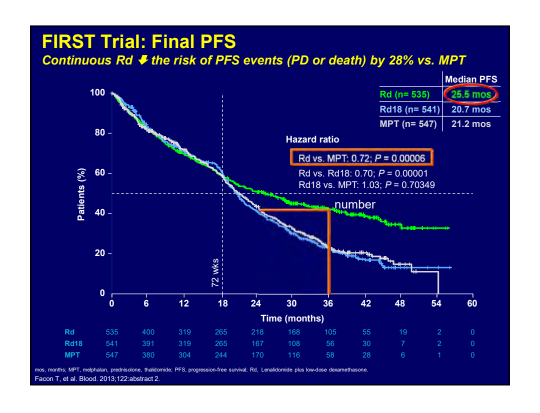


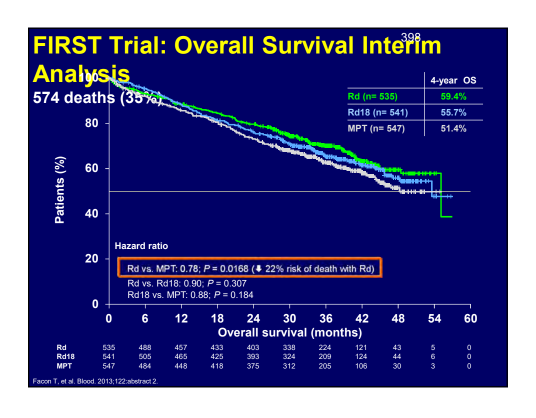




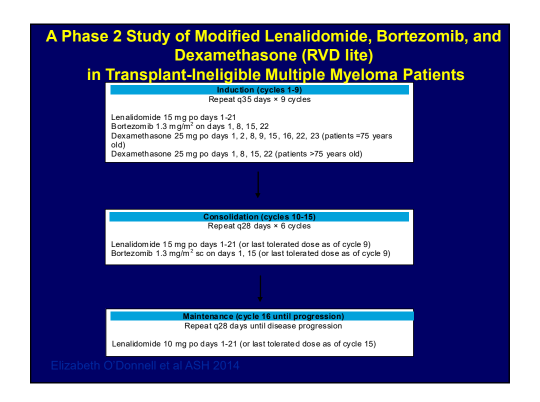




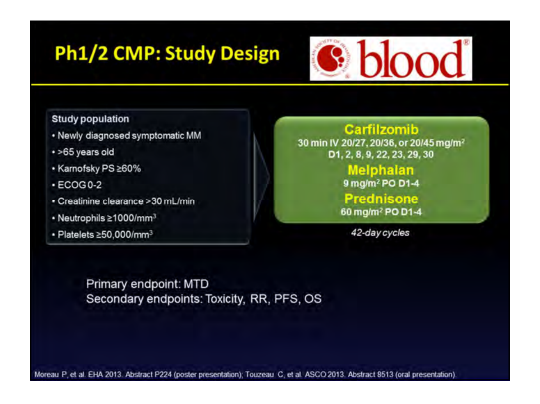


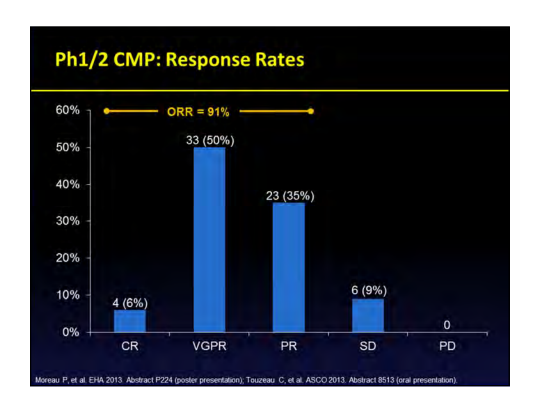


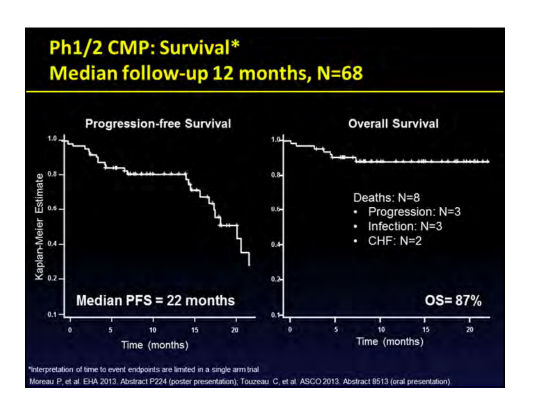
The Older Fit Patient Therapeutic Considerations: Is CR/MRD the goal? Is transplant the way? Is continued treatment best? Can novel drugs improve outcome?

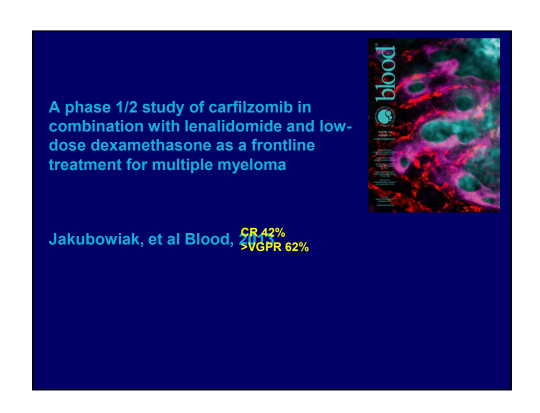


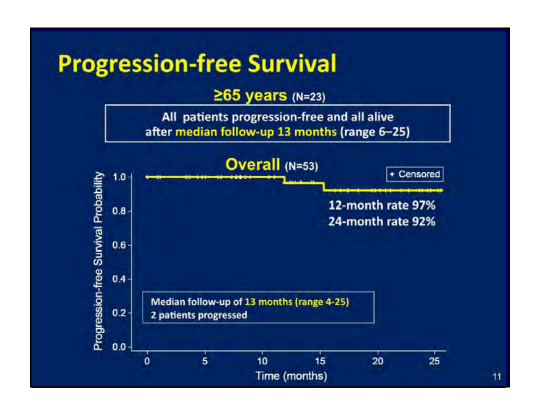
Response after	4 cycle	es (%) ((n=30)	
ORR (≥PR)	2	7 (90.0)	
CR		5 (16.7	7)	
VGPR		11 (36	7)	
PR		`	,	
FK		11 (36	.1)	
<u>SD</u>		3 (10.	0)	
VGPR or better	1	6 (53.3	5)	
Fatigue	9 (26.5)	7 (20.6)	1 (2.9)	17 (50.
Peripheral sensory neuropathy	7 (20.6)	6 (17.6)	1 (2.9)	14 (41.
Hypophosphatemia Edema limbs	1 (2.9)		11 (32.4)	12 (35.3
Rash maculo-papular	11 (32.4)		1 (2.9)	12 (35.3
Insomnia	3 (8.8)	4 (11.8)	4 (11.8)	11 (32.4
Depression	3 (8.8)	5 (14.7)	1 (2.9)	9 (26.
'	5 (14.7)	3 (8.8)		8 (23.
Diarrhea	7 (20.6)	1 (2.9)		8 (23.
Constipation	5 (14.7)	2 (5.9)		7 (20.
Dysgeusia	5 (14.7)	2 (5.9)		7 (20.
Hyperglycemia	4 (11.8)	2 (5.9)	1 (2.9)	7 (20.
Psychiatric disorders	2 (5.9)	3 (8.8)	2 (5.9)	7 (20.
Skin and subcutaneous tissue				6 (17.

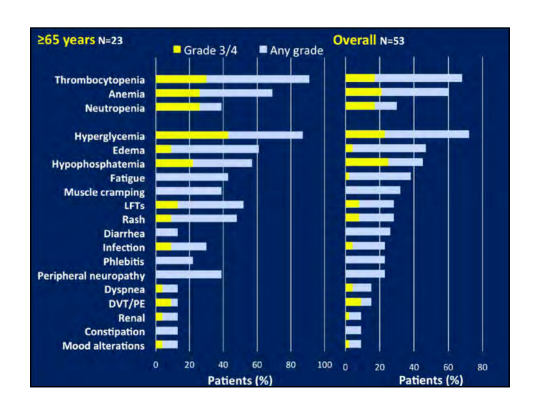


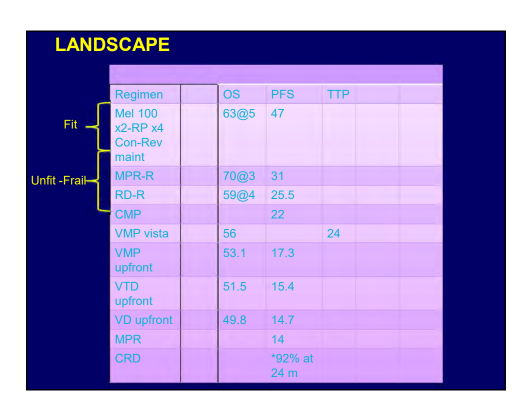


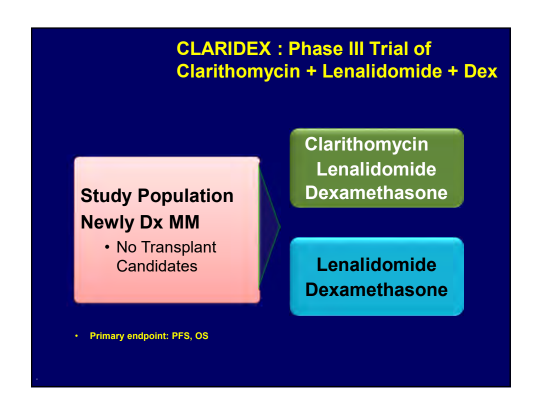


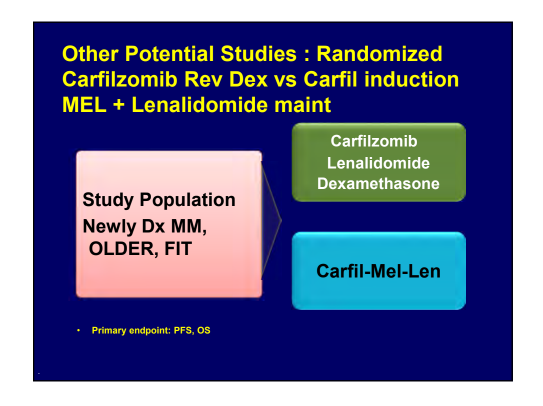


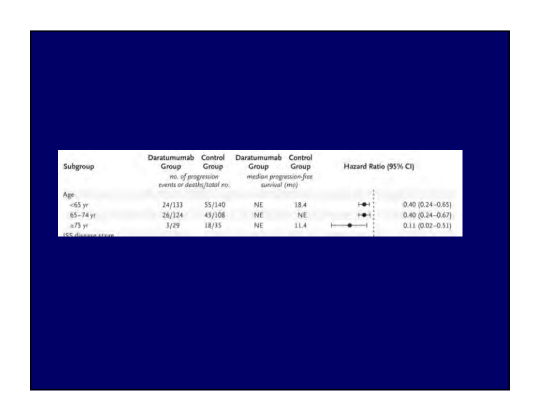






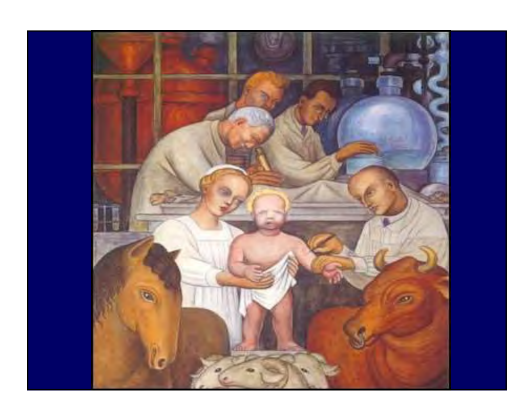






The Older Fit Patient

Therapeutic Considerations:
Is CR/MRD the goal? YES
Is transplant the way? Can be
Is continued treatment best? YES
Can novel drugs improve outcome? YES



Collaborators

Myelomacenter.org

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Linda Tegnestam Kathleen Pogonowski Stanley Goldsmith MD Joseph Lane MD Paul Christos



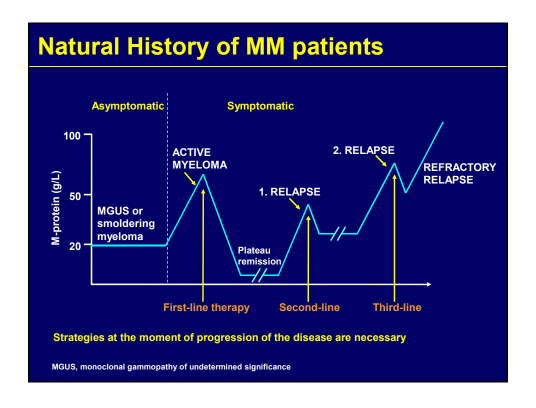


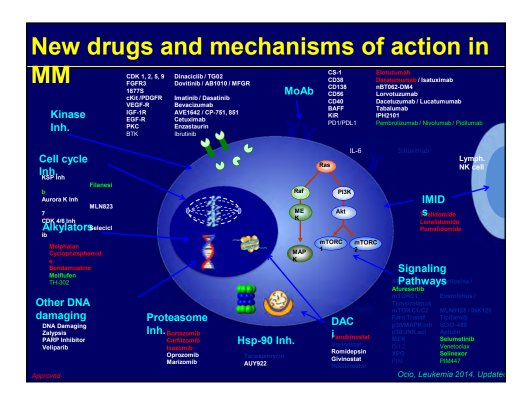


Cancer Research Center

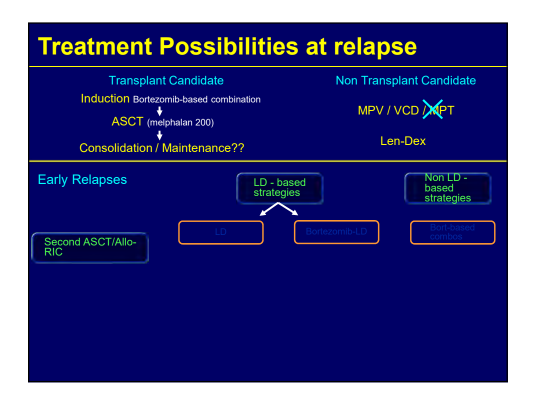
Tratamiento del MM en recaída

Enrique M. Ocio University Hospital & Cancer Research Center University of Salamanca Spain

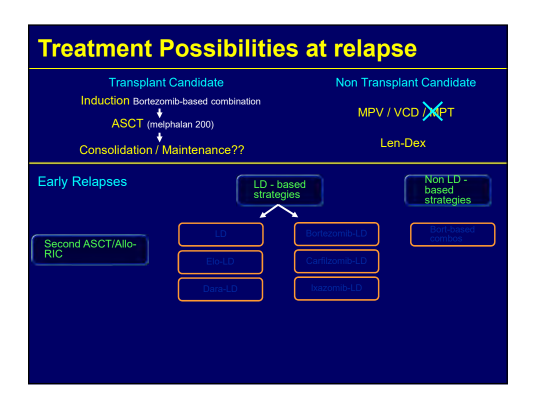


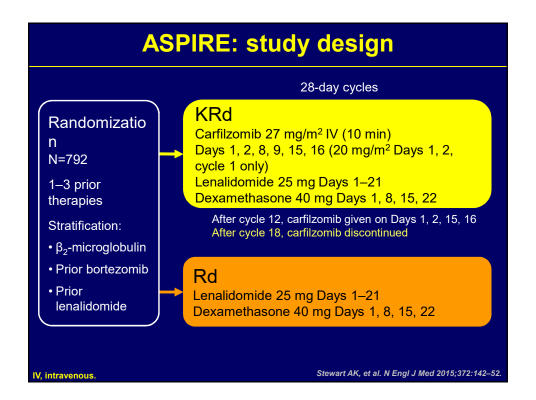


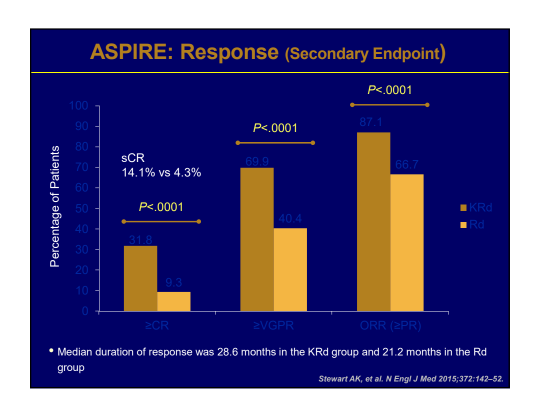
Which options do we have in early relapses?

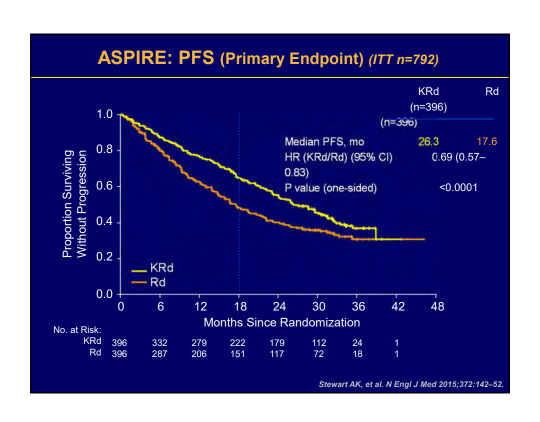


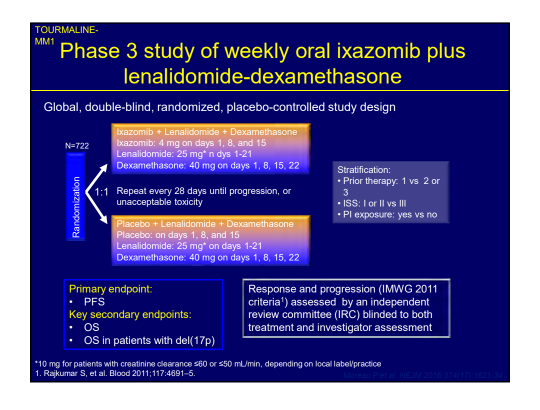
Is Len-Dex still a standard for early relapses?

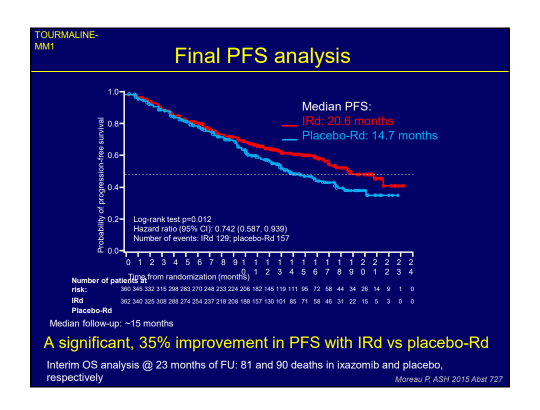








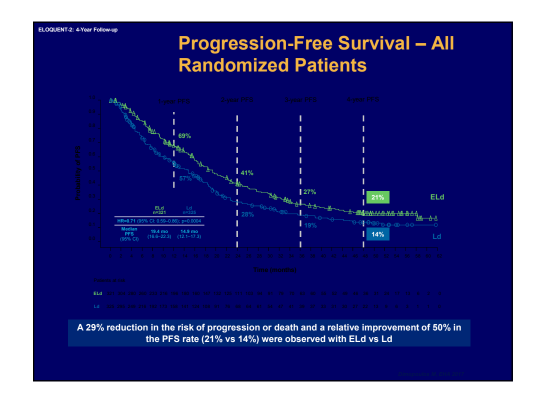




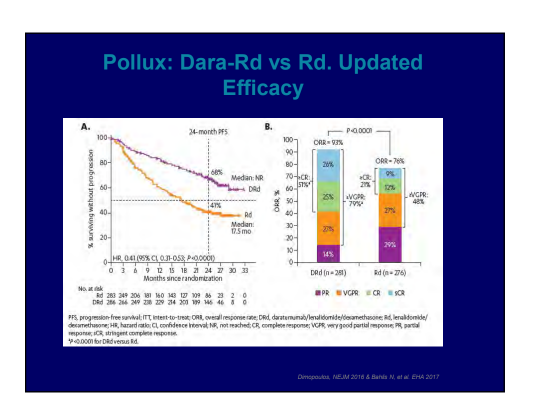
Eloquent-2: Elo + Ld vs Ld Study Design Open-label, randomized, multicenter, phase 3 trial (ELOQUENT-2) Elo plus Len/Dex (E-Ld) schedule (n=321) Key inclusion criteria Assessment Elo (10 mg/kg IV): Cycle 1 and 2: days 1, 8, 15, 22; Cycles 3+: days 1, 15 • Tumor response: RRMM 1–3 prior lines of therapy Prior Len exposure permitted in 10% of study every 4 wks until Len (25 mg PO): days 1-21 progressive disease, Dex: weekly equivalent, 40 mg Survival: every 12 wks after population (patients not refractory to Len) Len/Dex (Ld) schedule (n=325) Len (25 mg PO): days 1-21; progression Dex: 40 mg PO days 1, 8, 15, 22 Repeat every 28 days Endpoints:

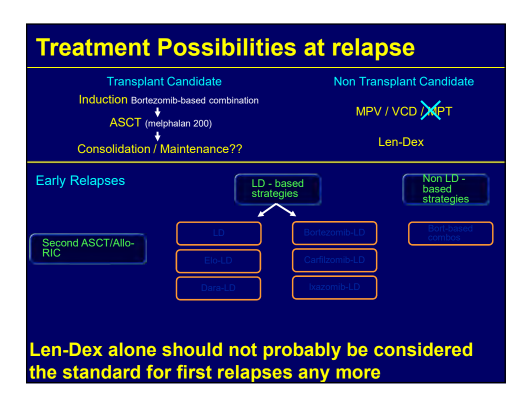


- Co-primary: PFS and ORR
- Other endpoints: overall survival (data not yet mature); duration of response, quality of life, safety
- All patients received premedication to mitigate infusion reactions prior to Elo administration

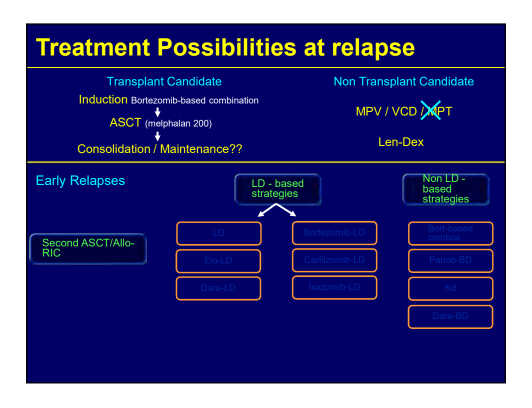


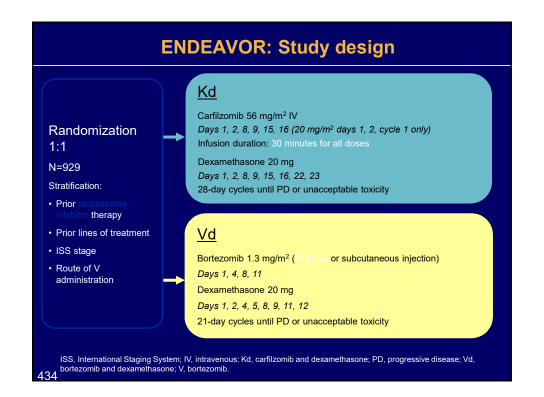
POLLUX: Study Design DRd (n = 286)Daratumumab 16 mg/kg IV Qw in Cycles 1-2, q2w in Cycles 3-6, then q4w until PD Key eligibility criteria Primary endpoint PFS A N RRMM R 25 mg PO Days 1-21 of each cycle until PD d 40 mg PO ≥1 prior line of D O Secondary endpoints therapy 40 mg weekly until PD • TTP Prior lenalidomide · os М exposure, but not Rd(n = 283)refractory • ORR, VGPR, CR Patients with MRD creatinine clearance Days 1-21 of each cycle until PD d 40 mg PO · Time to response ≥30 mL/min Duration of 40 mg weekly until PD Stratification factors Statistical analyses · No. prior lines of therapy • 295 PFS events: 85% power for Cycles: 28 days · ISS stage at study entry 7.7 month PFS improvement · Prior lenalidomide • Interim analysis: ~177 PFS events Pre-medication for the DRd treatment group consisted of dexamethasone 20 mga, paracetamol, and an *On daratumumab dosing days, dexamethasone was administered 20 mg premed on Day 1 and 20 mg on Day 2; RRMM, relapsed or refractory multiple myeloma; ISS, international staging system; R, lenalidomide; DRd, daratumumab/lenalidomide/dexamethasone; IV, intravenous; qw, once weekly; q2w, every 2 weeks; q4w, every 4 weeks; PD, progressive disease; PO, oral; d, dexamethasone; Rd, lenalidomide/dexamethasone; TTP, time to progression; MRD,

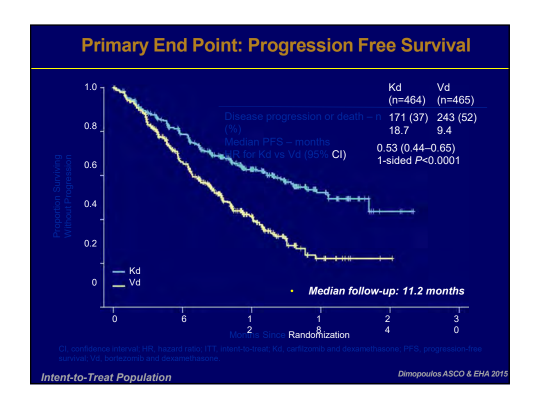


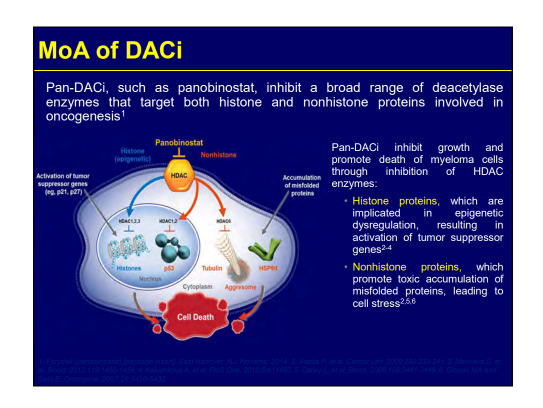


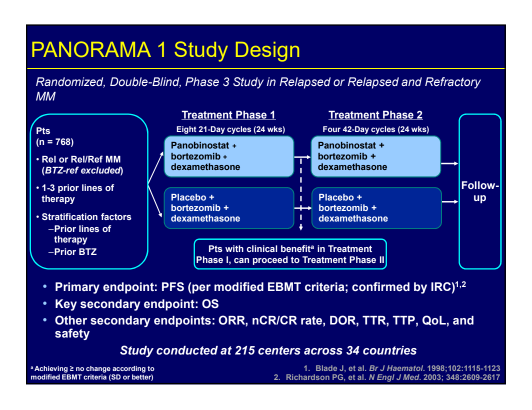
Which PI-based possibilities do we have?

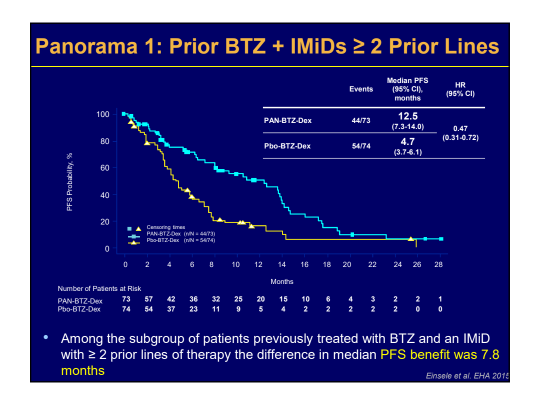












Panorama 1: Toxicity

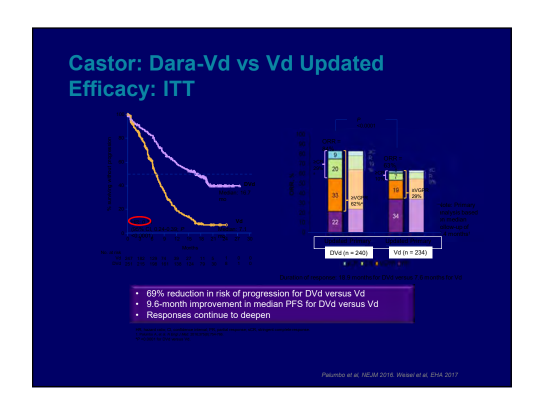
Median time on treatment of 152 days (3-411) for Panobinostat + Bort + Dex vs 187 days (3-443) for the control arm

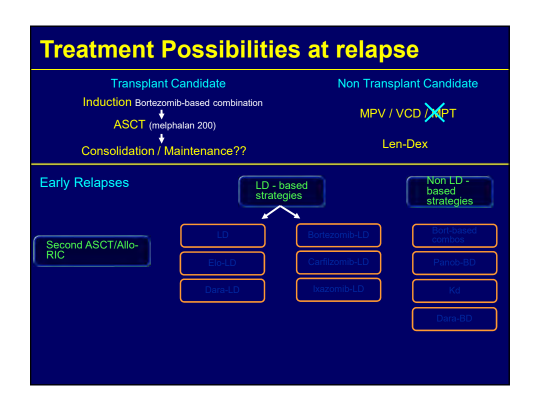
33% of pts discontinued treatment due to AEs (vs 17% in the control arm)

PAN-BTZ-D	ex (n = 381)	Pbo-BTZ-De	x (n = 377)
All grades	Grade 3/4	All grades	Grade 3/4
68.2	25.5	41.6	8.0
60.6	17.6	67.1	14.6
57.0	23.9	40.6	11.9
36.2	5.5	20.7	0.5
28.6	2.1	19.1	0.3
28.1	3.1	12.5	1.1
26.8	1.0	32.6	1.1
26.0	1.3	14.9	1.9
25.7	7.3	13.0	1.3
21.3	1.0	18.6	0
97.6	67.4	83.5	31.4 an Miguel, Lance
	All grades 68.2 60.6 57.0 36.2 28.6 28.1 26.8 26.0 25.7 21.3	68.2 25.5 60.6 17.6 57.0 23.9 36.2 5.5 28.6 2.1 28.1 3.1 26.8 1.0 26.0 1.3 25.7 7.3 21.3 1.0	All grades Grade 3/4 All grades 68.2 25.5 41.6 60.6 17.6 67.1 57.0 23.9 40.6 36.2 5.5 20.7 28.6 2.1 19.1 28.1 3.1 12.5 26.8 1.0 32.6 26.0 1.3 14.9 25.7 7.3 13.0 21.3 1.0 18.6 97.6 67.4 83.5

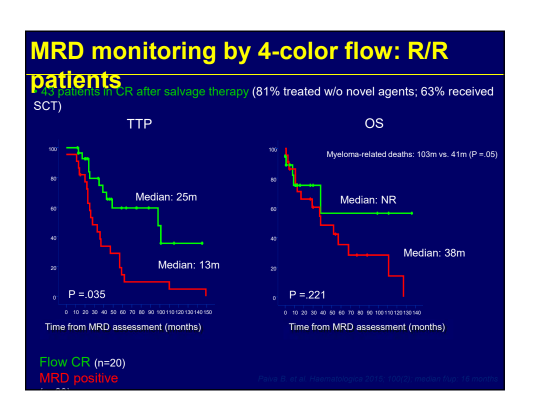
Oncology 2014

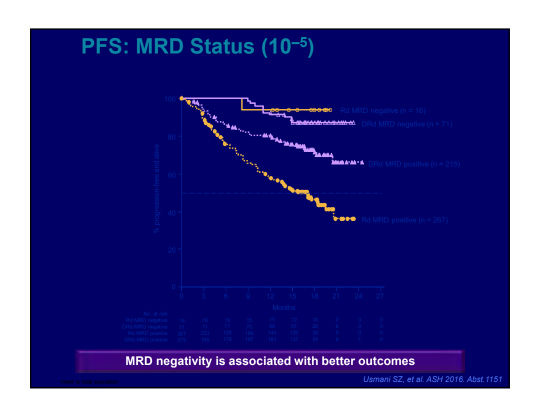
CASTOR: Study Design Multicenter, randomized, open-label, active-controlled phase 3 study DVd (n = 251)R Primary Endpoint Daratumumab (16 mg/kg IV) A N D O Every week - cycle 1-3 Every 3 weeks - cycle 4-8 Every 4 weeks - cycles 9+ Key eligibility criteria Secondary Endpoints Vel: 1.3 mg/m² SC, days 1,4,8,11 - cycle 1-8 dex: 20 mg PO-IV, days 1,2,4,5,8,9,11,12 - cycle 1-8 ORR, VGPR, CR М Prior bortezomib • MRD Vd (n = 247) Time to response refractory Z E · Duration of response Vel: 1.3 mg/m² SC, days 1,4,8,11 - cycle 1-8 dex: 20 mg PO-IV, days 1,2,4,5,8,9,11,12 - cycle 1-8 Cycles 1-8: repeat every 21 daysCycles 9+: repeat every 28 days Daratumumab IV administered in 1000 mL to 500 mL; gradual escalation from 50 mL to 200 mL/min permitted RRMM, relapsed or refractory multiple myeloma; DVd, daratumumab/bortezomib/dexamethasone; IV, intravenous; Vel, bortezomib; SC, subcutaneous; dex. dexamethasone; PO, oral; Vd, bortezomib/dexamethasone; PFS, progression-free survival; TTP, time to progression, ORR, overall response rate; VGPR, very good partial response; CR, complete response; MRD, minimal residual disease.

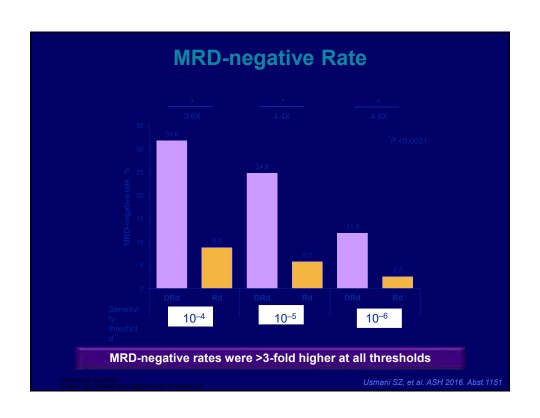


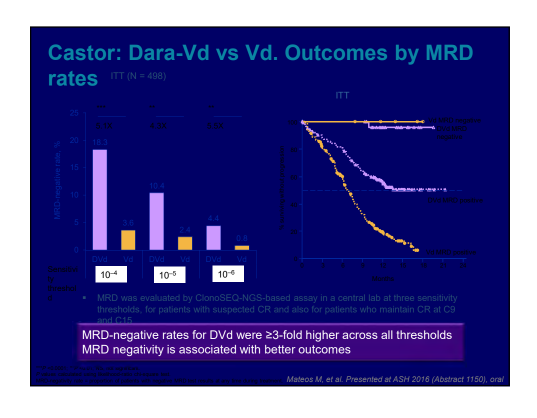


Is it possible and important to achieve MRD- in RRMM?

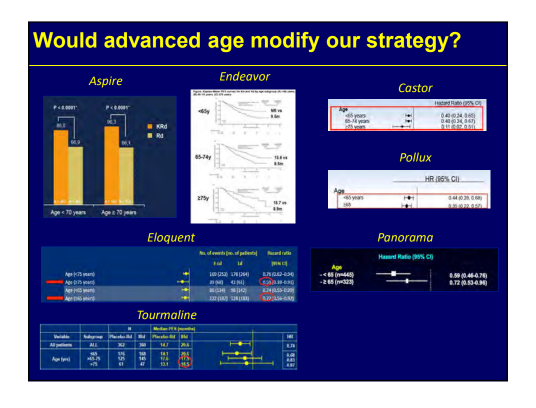






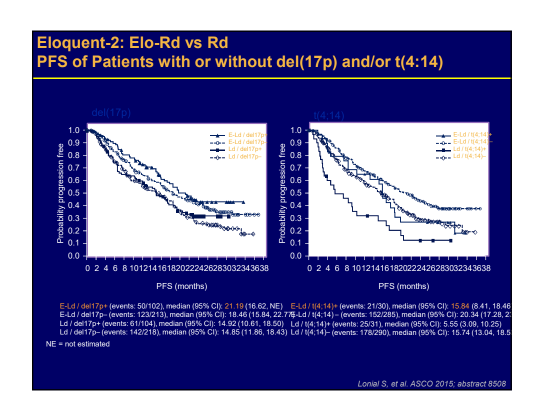


Would advanced age modify our strategy?



What if the patient has adverse cytogenetics?

ASPIRE: KRd vs Rd in RMM Subgroup analysis in HR patients: PFS High-risk group Standard-risk group KRd Rd KRd Rd (n=48)(n=52)(n=147)(n=170)PFS, PFS. median 13.9 median 19.5 months months Hazard Hazard 0.70 0.66 ratio (95% ratio (95% (0.43 - 1.16)(0.48 - 0.90)CI) CI) Avet Loiseau H, ASH 2015 Abst



Tourmaline-MM1: I-Rd vs Placebo-Rd in RMM

Subgroup analysis in HR patients: PFS

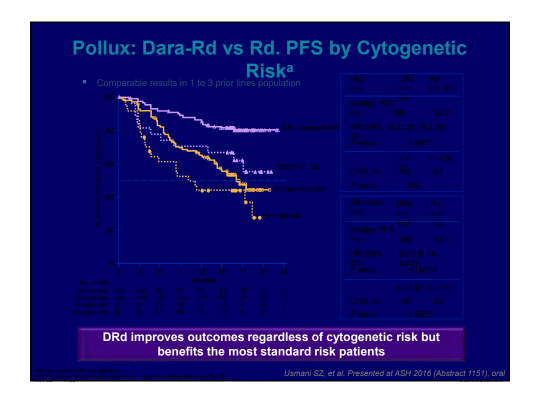
			≥VGPR, %		≥CR, %				
			IRd		IRd		IRd		
All patients	78.3 *	71.5	48.1 *	39	11.7	6.6	20.6	14.7	0.742
Standard-risk patients	80	73	51	44	12	7	20.6	15.6	0.640
All high-risk patients	79*	60	45*	21	12*	2	21.4	9.7	0.543
Patients with del(17p) [†]	72	48	39	15	11*	0	21.4	9.7	0.596
Patients with t(4;14) alone	89	76	53	28	14	4	18.5	12.0	0.645

*p<0.05 for comparison between regimens. †Alone or in combination with t(4;14 or t(14;16)

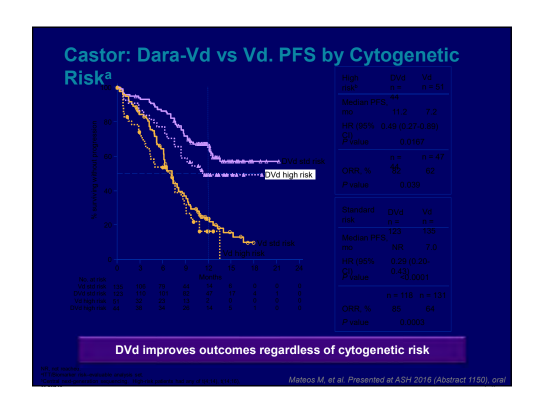
- Median OS was not reached in either arm
- In the IRd arm, median PFS in high-risk patients was similar to that in the overall patient population and in patients with standard-risk cytogenetics

Cut-off values: del(17p) >5%: t(4:14) and t(14:16) >3%

Moreau P, ASH 2015 Abst 727



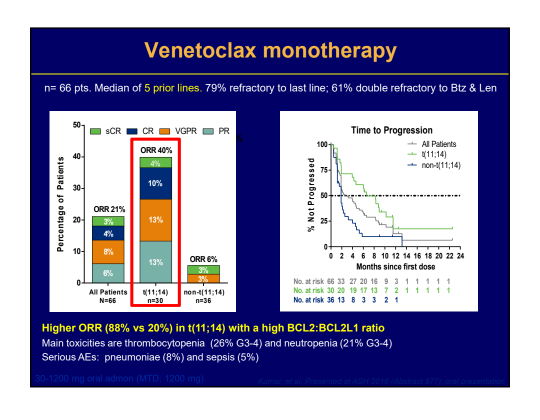
Endeavor: Carfilzomib-Dex vs Bortezomib-Dex in RMM Subgroup analysis in HR patients: PFS High-risk group Standard-risk group Kd Vd Kd Vd (n=284) (n=97) (n=113) (n=291) PFS, median PFS, median 6.0 10.2 (6.9–11.3) (18.7-NE) months (4.9 - 8.1)months (9.3-12.2)(95% CI) (95% CI) Hazard Hazard 0.646 (0.453–0.921) 0.439 (0.333–0.578) ratio (95% ratio (95% CI) CI) Chng WJ, ASH-2015 Abst 30

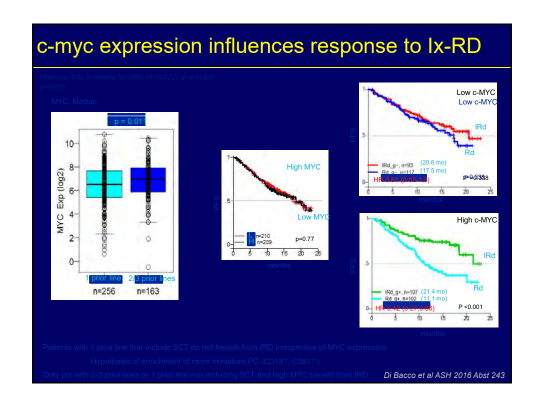


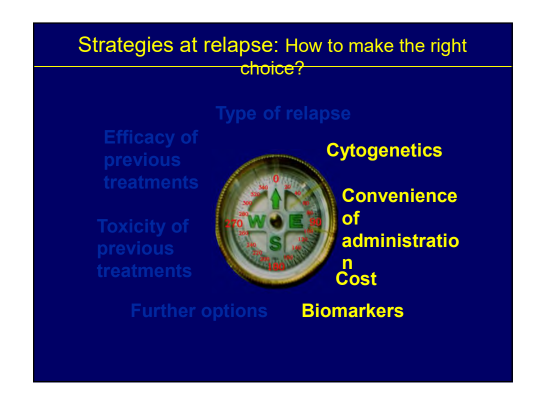
Efficacy of novel combinations based on cytogenetic risk

	Exp	Contro	gh Risk Dif.	HR	Exp	Contro	St Risk Dif.	HR
	EXP	+ + -	DII.	пк	EXP	-	DII.	nk
Aspire ¹ K-Rd vs Rd	23.1	13.9	9.2	0.70 (0.43–1.16)	29.6	19.5	10.1	0.66 (0.48–0.90)
Tourmaline ² IRd vs Rd	21.4	9.7	11.7	0,543	20.6	15.6	5	0.640
Eloquent 2 ³ Elo-Rd vs Rd	21.19	14.92	6.27	-	18.46	14.85	3.61	-
Eloquent 2 ³ Elo-Rd vs Rd t(4;14)	15.84	5.55	10.29	-	20.34	15.74	4.6	-
Pollux ⁴ Dara-Rd vs Rd	NR	10.2	-	0.44 (0.19-1.03)	NR	17.1	-	0.30 (0.18-0.49
Endeavor⁵ Kd vs Bd	8.8	6	2,8	0.646 (0.453-0.921)	NE	10.2	-	0.439 (0.333-0.578)
Castor ⁶ Dara-Bd vs Bd	11.2	7.2	5	0.49 (0.27-0.89)	NR	7.0		0,29 oiseau.H. ASH 2 u.P. ASH 2015 A

Can we use biomarkers to predict sensitivity to a given combination?

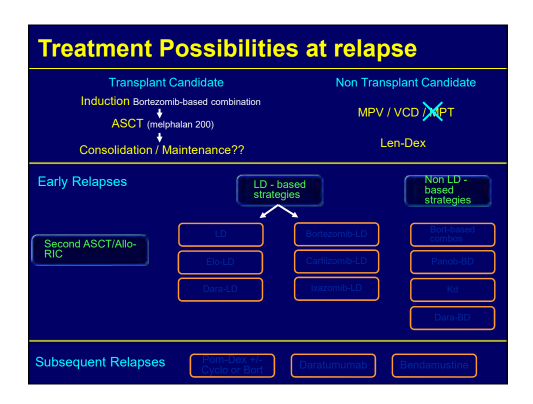






Strategies at relapse: How to make the right choice? 1. Does this patient require treatment? 2. Is he candidate to ASCT at relapse 3. Which were the prior regimens: Efficacy and toxicity? Candidate to Len-Dex based Candidate to Non Len-Dex based K-Rd I-Rd Bd retr. Panob-Bd Elo-Rd Dara-Rd Dara-Bd 4. Factors to take into considerations ive rel. (more potent) ... K-Rd; Dara- • Indolent relapses (>3.5 y TFI) ... Elo-Rd 1 prior line ... K-Rd; Dara-Bd; Kd Bd Prior PI refr. ... Elo-RD; Dara-Rd; к-Rd, I-Rd • Preference for oral ... I-Rd High risk cytog. ... K-Rd; I-Rd; Elo-Rd; Dara-Vd, Dara

Which options do we have in late relapses?



➤ Single agent¹ (31 patients relapsing HDT) ➤ Benda-Bort² (40 patients 6 prior lines) ORR: 27% (2% CR, 5% VGPR, 21%PR) ► Benda-Bortz-Dex³ (40 patients 4 prior lines) ORR: 72% (25% VGPR, 47%PR) ➤ Benda-Bort-Dex⁴ (79 patients 2 prior lines) ORR: 61% (15% CR, 20% VGPR, 25% PR) ► Benda-Bort-Dex⁵ (73 patients elderly 1st rel.) ORR: 70% (14% CR, 16% VGPR, 40% PR) ➤ Benda-Bort-Dex⁶ (75 patients 1 prior line) ORR: 72% (16% CR, 19% VGPR, 37% PR) ➤ Benda-Bort-Pred⁷ (78 patients 2 prior lines) ➤ Benda-Thal-Pred⁸ (28 patients) ➤ Benda-Thal-Dex⁹ (23 patients ➤ Benda-Thal-Dex¹⁰ (66 patients ➤ Benda-Len-Dex¹¹ (29 patients 3 prior lines) ➤ Benda-Len-Dex¹² (41 patients 3 prior lines) ORR: 50% (11% CR, 7% VGPR, 32% PR) 1. Knop et al. Hematologica 2005, 5. Rodon, ASH 2013. Abstract 1971 9. Grey-Davies E. BJH 2012 90:8287son. BJH 2013

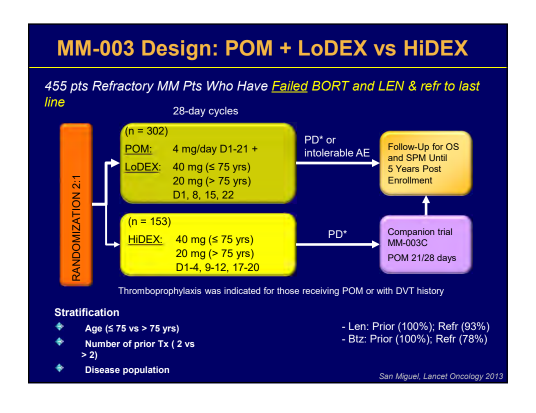
20B2Rnson. BJH 2013 6. Offidani, Blood Cancer J 2013 10. Schey, ASH 2013. Abstra 3. Hrusowsky et al ASH 2007 Abstract 4851 7. Pönisch et al. J Cancer Res Clin Oncol 2013 11. Lentzsch. S. Blood 2012

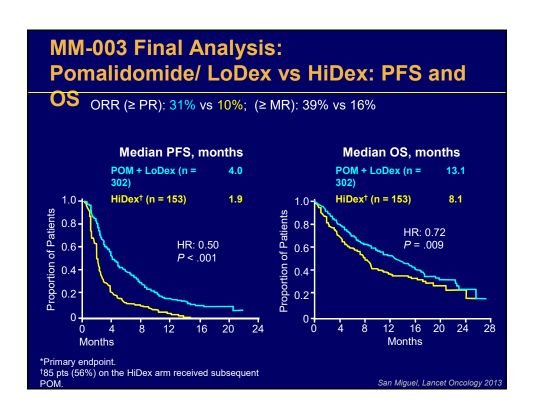
8. Pönisch et al. BJH 2008,

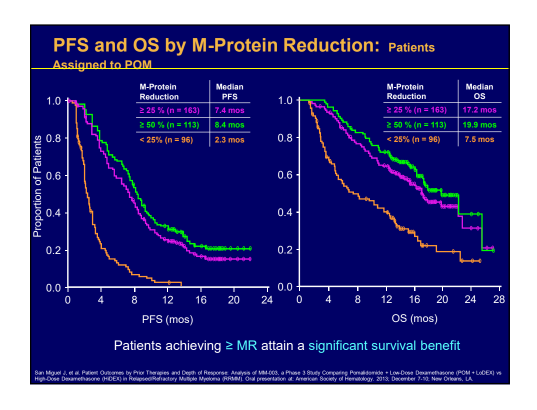
12. Pozzi, ASH 2013. Abstract 3222

Bendamustine in R/R MM

4. Ludwig H. Blood 2013

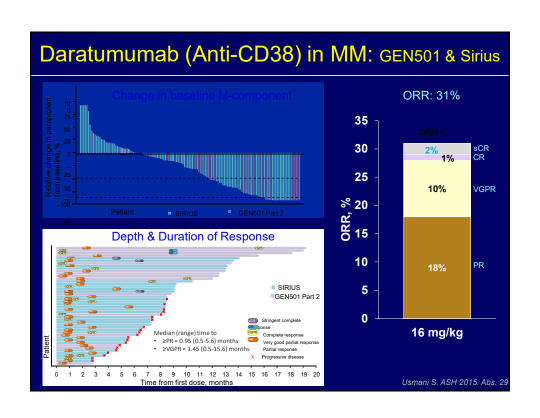




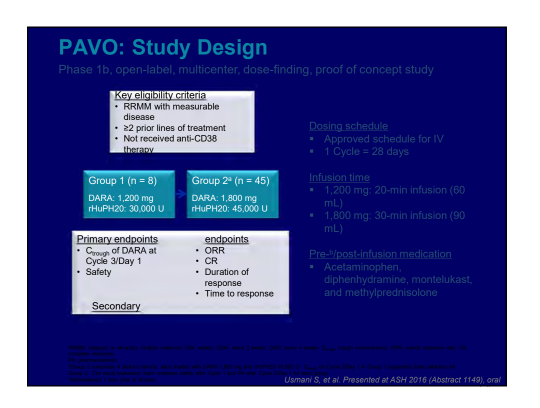


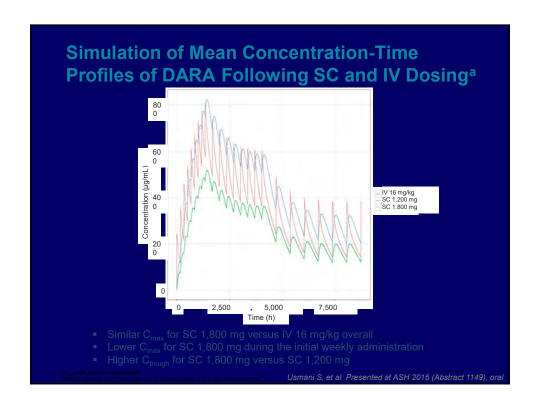
	POM + Vd ¹	K + POMdex ²	Ixa + POMdex³	Dara + POMdex ⁴	
Study phase	1	1/2	1/2	1	
Prior lines of therapy, n	1–4		1–5 including PI and Len	≥2 (2–13)	
Refractory to Len, n (%)	All patients were Len- refractory	40 (87)	32 (100); 25 (100)	87 (89)	
Refractory to PI, n (%)	All pts were PI-exposed (but not refractory)	NR	20 (63); 15 (60)*	74 (76)	
ORR, %	65	84	44	71	
Median (range) DOR *bortezomib + MTD: POM Median PFS months QZW+POM 4 mg D1-21	7.4 (4.4–9.6) months 4 mg + BORT (IV or SC) 1.3 m + Dex 40/20 mg QW	NR g/m² + LoDex 20 mg (10 mg f 12.9	56 (28-160) months or patients >75 years); ‡MT	NR D: Carfilzomib 27 mg/m² 6-month rate = 66%	
D, day; Dex, dexamethasone; DO progression-free survival	R, duration of response; IMiD, Immunomod				
• Pomalidom	ide combinations at	ASH 2016			
	Marizomib; Carfilzomib; ; Selinexor; ACY-247	Ixazomib; Daratumuma	ab; MOR202; Isatuxii	nab; Pembrolizumab;	
	et al. ASH 2016. Absi Spencer et al,	ASH 2016 Abs. 332@ringhen et	al, ASH 2016 Abs. 114 5 Kumar	et al, ASH 2016 Abs. 3327	

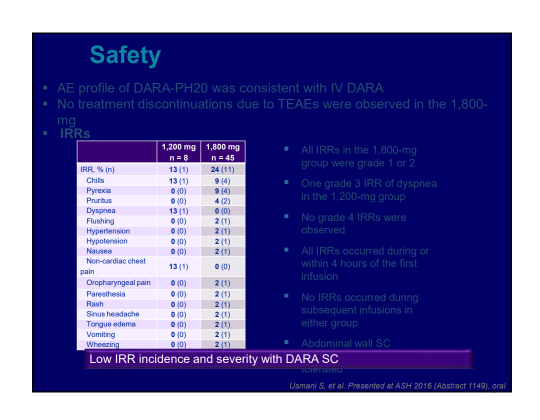
Daratumumab (Anti-CD38): Background Daratumumab binds to CD38 Human CD38 IgGk monoclonal Direct ON-TUMOR Actions antibody **IMMUNOMODULATORY** Actions Modulation of tumor Direct and indirect antimyeloma activity1-5 Depletes CD38+ immunosuppres sive regulatory Promotes T-cell expansion activation5 MYELOMA CELL DEATH

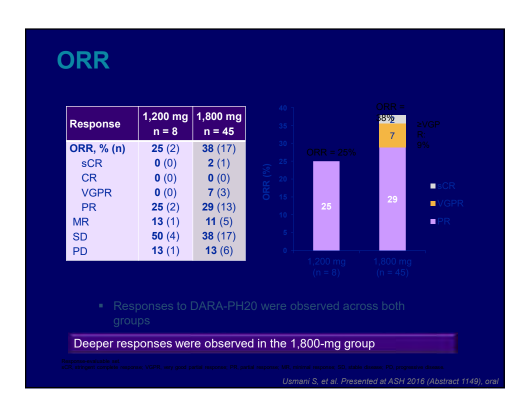


Progression-free survival Overall Survival Responders MR/SD: 17.5 (15.1-NE) PDINE: 0.9 (0.9-1.0) months PDINE: 0.9 (0.9-1.0) months PDINE: 0.9 (0.9-1.0) months PDINE: 0.9 (0.9-1.0) months Responders MR/SD: 17.5 (15.1-NE) PDINE: 0.9 (0.9-1.0) months Responders MR/SD: 17.5 (15.1-NE) PDINE: 0.9 (0.9-1.0) months PDINE: 0.9 (0.9-1.0) months PDINE: 0.9 (0.9-1.0) months Responders MR/SD: 17.5 (15.1-NE) Policetis at IRIN Responders MR/SD: 17.5 (15.1-NE) Policetis at IRIN Responders MR/SD: 17.5 (15.1-NE) Time from first does, months 1-year overall survival rate = 69% (95% CI, 60.4-75.6) Usmani S. ASH 2015. Abs. 29









Conclusions

- Identify Biochemical relapses → Does this patient require treatment?
- Early relapses (1-3 prior):
 - Len-Dex substituted by K-Rd; Elo-Rd; I-Rd; Dara-Rd;
 - · Non Len-Dex combinations: Kd; Panob-Bd and Dara-Bd
- Late relapses (>2-3): Bendamustine, Pom-Dex; Daratumumat
- Probably the problem has changed
 - What else can I prescribe to this patient? → How should I choose?
 - Based on: Efficacy & Tox of prev. Tx; Type of Relapse; Age; Cost; Cvtogenetics
- · Continue including patients in clinical trials so in the next 10 years





Salamanca



Tratamiento del MM en recaída

Enrique M. Ocio University Hospital & Cancer Research Center University of Salamanca Spain

Amyloidosis: "Under-diagnosed disorder"

Joan Bladé Amyloidosis and Myeloma Unit Hospital Clinic of Barcelona

Santiago de Chile, 12 de Agosto, 2017

Concept of Amyloidosis

 Group of "rare" diseases characterized by extracellular deposition of amyloid fibrils (Congo red +) in organs and tissues

Pathogenesis

Increased synthesis, specific mutations or aging of autologous proteins (amyloidogenic precursor proteins)

Conformational changes and aggregation forming amyloid fibrils (Congo-red positive)

Deposition of amyloid fibrils in tissues cause functional damage of involved organs, and eventually (if untreated) leads to death

Types of Amyloidosis

Can be classified according to:

- Localized versus systemic
- Acquired versus hereditary
- Different amyloidogenic precursor proteins

Most Common Forms of Systemic Amyloidosis 99%

Precursor protein	Main synthesizing organ	Amyloid type
Ig Light chain	Bone marrow	Ig LC amyloidosis 75%
Serum amyloid A protein	Liver	Secondary (reactive) amyloidosis = AA amyloidosis
Wild type transthyretin	Liver	Senile systemic amyloidosis (SSA) = wild type ATTR amyloidosis
Mutated transthyretin	Liver	Hereditary ATTR amyloidosis = hereditary ATTRV30M amyl.
Mutated apolipoprotein A1	Liver, GI tract	Hereditary AApoAI amyloidosis

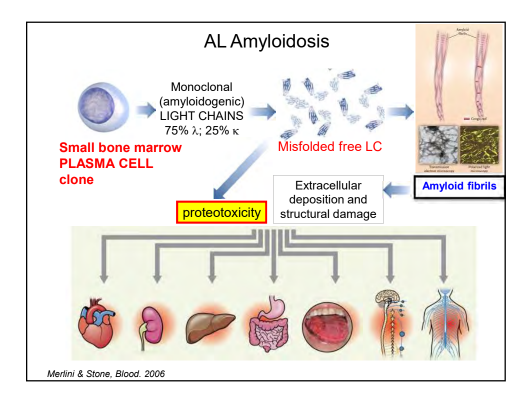
Localized Amyloidosis

- Local formation and deposition of light chain (AL) amyloid fibrils, confined to a single site (WITHOUT an underlying systemic plasma cell disorder)
- Sites:
 - tracheo-bronchial tree (upper resp tract, nasopharynx),
 - urinary tract (bladder),
 - skin and nails,
 - GI tract,
 - Others (lymph nodes, orbit, ...)
- Treatment: LOCAL, if needed (surgery, laser, RDT...)
- Low risk of progression to systemic AL amyloidosis

Light Chain Amyloidosis (AL Amyloidosis)

Epidemiology

- Incidence: 4-10 patients/million/year
- Aprox. one-fifth as common as multiple myeloma
- Median age at diagnosis: 64 years
 (<5% of patients are < 40 years-old)
- Male predominance: 2/3 of patients (referral bias?)
- Median survival (untreated patients): 10-14 months



Monoclonal Gammopathy

- M protein by electrophoresis (moderate size): 50%
- M protein by immunofixation: 90-95%
- Isotype: 32% IgG
 24% Bence-Jones
 10% IgA,
 5% IgM
 1% IgD
- Light chain isotype: LAMBDA (75%)
- Bone marrow plasma cell infiltration: 5-10%
 (>20% in about ¼ patients)
- No CRAB

Main clinical manifestations

- Fatigue and anorexia (weight loss)
- Organ involvement
 - Kidney, 70-80%
 - Heart, 50-70%
 - Liver, 20%
 - Peripheral / autonomic neuropathy, 20%/15%
 - Gastrointestinal, 8% (only 1% symptomatic)
 - Soft tissue, 12%

Renal involvement (70-80%)

- Glomerular proteinuria evolving to overt nephrotic syndrome (hypercholesterolemia, edema)
- Less frequently renal failure progressing to ESRD
 - Serum creatinine > 2 mg/dL in 20% patients

Cardiac involvement (50-70%)

- Restrictive cardiomyopathy
- Congestive heart failure
- Arrhythmias (syncope or sudden death)
- Rarely angina or infarction
- Elevation of cardiac biomarkers (NT-proBNP, troponin T or I)

Cardiac involvement (50-70%)

- ECG: low voltages +/- pseudoinfarct pattern
- Echo: LV concentric hypertrophy
 - Thickening of LV and RV walls, interventricular and interatrial septa (>12 mm)
 - "Granular sparkling"
 - Normal LV ejection fraction (reduced in advanced disease)

Cardiac MR: late gadolinium ennancement

Liver involvement (20%)

- Hepatomegaly
 - (Amyloid infiltration versus congestive heart failure in patients with cardiac amyloidosis)
- Elevated alkaline phosphatase
- Rarely elevated bilirubin
 - (bilirubin > 5 mg/dL ⇒ survival <1 month)
- Kappa clones more frequently found (30-40%)

Nervous system involvement

Peripheral neuropathy (PN) (20%)

- Symmetric distal sensori(motor) PN
- Seldom isolated (2%) in contrast with PN in ATTR-V30M

Autonomic neuropathy (15%)

- Postural hypotension
- Impotence
- Bowel dysfunction (severe diarrhea to constipation)

GI involvement

- Bleeding (due to vascular fragility and loss of vasomotor responses to injury)
- Gastroparesis
- Malabsorption
- Constipation
- Intestinal pseudo-obstruction (dysmotility)

Other clinical manifestations (I)

Periorbital purpura (15%)



- Macroglossia (10%)
- Submandibular swelling
- Hoarse or weak voice
- Dry mouth
- Jaw claudication



Other clinical manifestations (II)

- Carpal tunnel syndrome (25%)
- Factor X deficiency (10-15%)
- Muscular pseudohypertrophy, articular deposits, shoulder pad sign
- Splenomegaly (5%)
 - Hyposplenism in 25% of patients
- Pulmonary/pleural involvement (<5%)

Diagnosis

- 1. Clinical suspicion:
 - ⇒ Any clinical manifestation
 - ⇒ Serum and/or urine M component
- 2. Confirmatory biopsy:
 - ⇒ Amyloid deposition (Congo Red +)
 - ⇒ Typing by immunohistochemistry (k or λ LC)
- 3. Gold-standard of congophilic deposits typing: Proteomics (LMD/MS)

Cohen AD, Comenzo RL. ASH 2010.

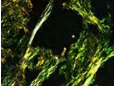
Serum Free Light Chain (FLC)

- Ig FLC is the precursor of amyloid
- FLC assay (Freelite)
- Role in diagnosis: Abnormal FLC in 90% of patients
- Role in disease monitoring:
 - "Measurable" by M-protein in 25% of patients
 - "Measurable" by FLC in >50% of patients
- Independent prognostic value

Proving systemic amyloid deposition

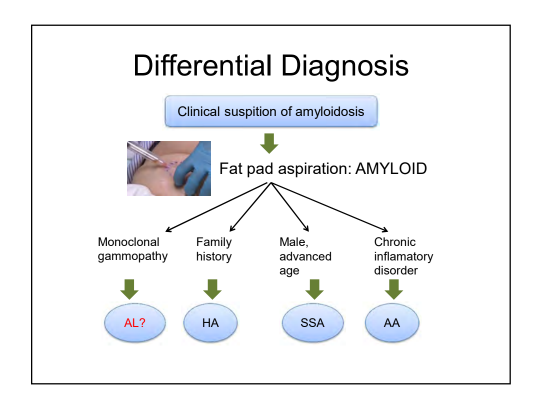
Diagnosis of amyloidosis relies on Congo red staining of tissue biopsy





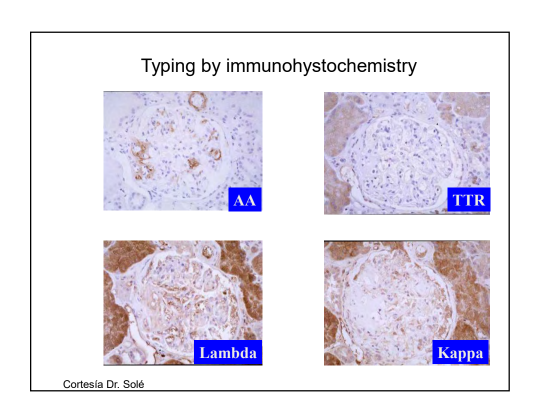
- Tissue of choice: abdominal fat (sensitivity 88%)
- If negative → labial minor salivary glands (S: 58%) or rectum
- If negative → involved organ (kidney, liver, heart,...) (hemorrhagic risk!!)

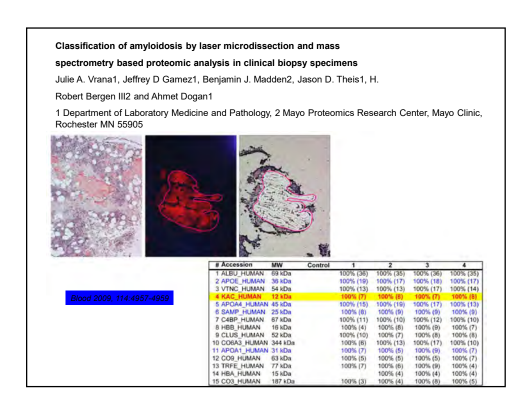
Courtesy of Dr. Merlini

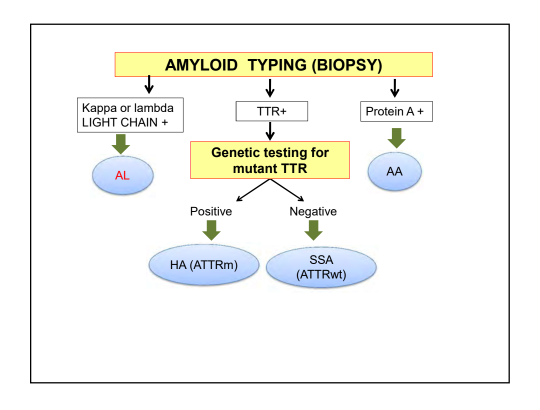


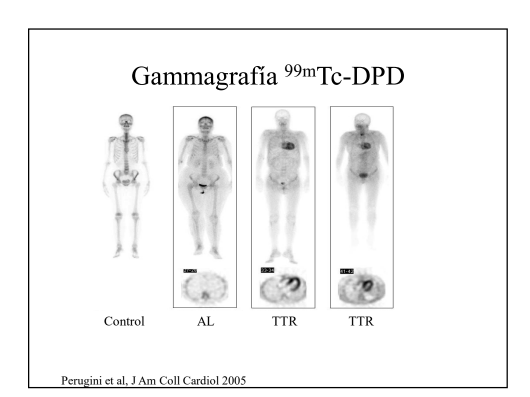
Treatment Approaches in Systemic Amyloidosis

Amyloid type	Treatment options
AL amyloidosis	Chemotherapy
AA amyloidosis	Treatment of underlying disease
Senile systemic amyloidosis (SSA)	Treatment of heart failure, heart Tx
Hereditary ATTR amyloidosis	Liver Tx
Hereditary AApoAI amyloidosis	Liver Tx









Looking for Organ Involvement

- Thorax X-ray
- ECG and echocardiogram
- Holter (suspected arrhythmia)
- Cardiac MR (unclear echo)
- Skeletal survey (bone pain)
- Electrophysiological studies (suspected polyneuropathy)
- GI endoscopic study (suspected GI involvement)

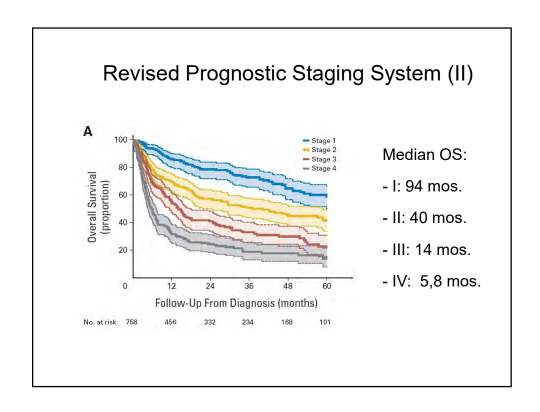
AL Prognosis

- Cardiac involvement → cardiac biomarkers
- Tumor burden → serum free light chains
- Response to therapy

Revised Prognostic Staging System (I)

Kumar et al, JCO 2012 (Mayo Clinic)

- N: 810 newly AL
- Prognostic factors for OS
 - FLC-diff ≥ 180 mg/L
 - -cTnT ≥ 0.025 ng/mL
 - NT-proBNP ≥ 1800 pg/mL



Hem Response (HR) Evaluation

	Gertz et al, 2005
CR	Negative serum and urinary immunofixation Normal FLC ratio BMPC <5%
PR	If serum M protein >5 g/L \rightarrow 50% \downarrow If urinary M protein > 100 mg/day \rightarrow 50% \downarrow If iFLC > 100 mg/L \rightarrow 50% \downarrow
SD	No CR, PR or progression

	Palladini et al, 2012
aCR	= ¿BM needed?
VGPR	dFLC* <40 mg/L
PR	dFLC* decrease >50%
SD	No response

^{*} Baseline dFLC must be ≥50 mg/L

Cardiac Response Evaluation

	Gertz et al, 2005	Palladini et al, 2012
Response	↓ IVSd (≥ 2 mm) or ↑ EF (20%) or ↓ NYHA (2 classes) No ↑ in diuretic use No ↑ in wall thickness	NT-proBNP* ↓ >30% and >300 ng/L NYHA class response (2 classes)
Progression	↑ IVSd (2 mm) ↑ NYHA (1 grade) with a ↓ EF of ≥ 10%	NT-proBNP* ↑ >30% and >300 ng/L cTn (I or T) ↑ >33% EF ↓ by 10% or more

^{*} If baseline level of NT-proBNP > 650 ng/L

Important: prognostic value of NT-proBNP at 3 and 6 months; prognostic value of cTn only at 6 months after start of therapy

Renal Response Evaluation

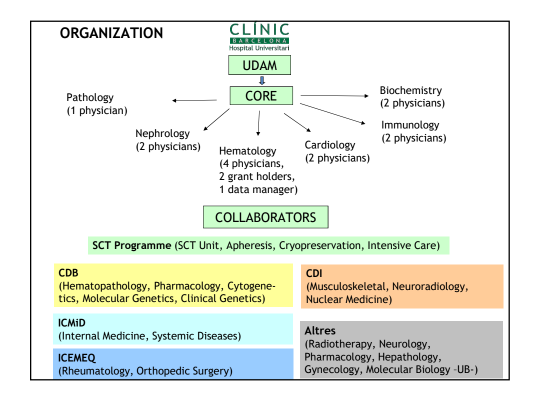
	Gertz et al, 2005	Palladini et al, 2014** (at 6 months)
Response	 -≥ 50% ↓ (at least 0.5 g/day) of 24-hr urine protein* Serum creat and creatCl must not worsen by 25% over baseline 	 ≥ 30% ↓ of 24-hr urine protein or below 0.5 g/24h No renal progression
Progression	 - ≥ 50% ↑ of 24-hr urine protein to at least 1g/day - or ≥ 25% worsening of serum creat or creatCl 	– ≥ 25% worsening of eGFR

^{*} If baseline level 24-hr urine protein > 0.5 g/day

^{**} Palladini et al, Blood 2014;124:2325-32. N= 461 (validation cohort: 271).

Take-Home Message

- Most important: early diagnosis
 - early initiation of the most effective therapy









Institute of Biomedical Research of Salamanca

University of Salamanca

Cancer Research Center

Macroglobulinemia de Waldenström

Enrique M. Ocio University Hospital & Cancer Research Center **University of Salamanca** Spain

Gammapatías monoclonales

- Enfermedades de Células Plasmáticas
 - Variantes de Mieloma
- ariantes de mieroma

 Mieloma Sintomático

 Mieloma Sintomático variante con Amiloidosis

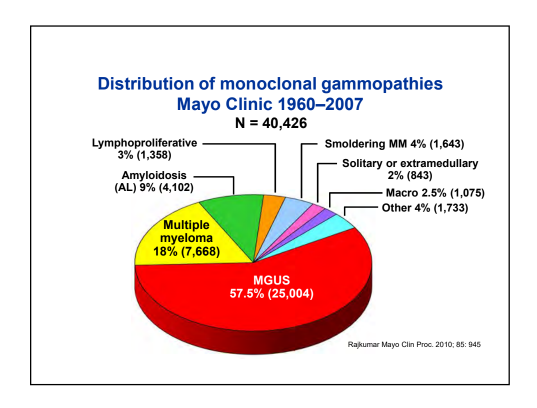
 Mieloma Sintomático variante con Enfermedad de Cadenas Ugeras Sistémica

 Mieloma Indolente ("Indolent")

 Mieloma Quiescente ("Smoldering")

 - Mieloma Osteosclerótico (Sindrome POEMS)
 Leucemia de Células Plasmáticas

 - Gammapatia Monoclonal de Significado Incierto (GMSI)
 - Plasmocitomas
 - Plasmocitoma Solitario del Hueso
 Plasmocitoma Extramedular
- Otras Enfermedades Inmunosecretoras
 - Macroglobulinemia de Waldenström (Inmunocitoma)
 - Enfermedad de Cadenas Pesadas (ECP)
 - · ECP gamma
 - ECP alfa
 - ECP alfa con enfermedad inmunoproliferativa del intestino delgado
 - Enfermedades de depósito de Inmunoglobulinas
 - · Enfermedad de Cadenas Ligeras Sistémica
 - Amiloidosis Primaria



Macroglobulinemia de Waldenström

Baja frecuencia

- 6% of todas la gammapatías monoclonales
- 2-5 casos/mill/año (GEM, 3'1) 2500 casos/año en Europa

Edad avanzada

· Mediana: 71. Masculino/Femenino: 2:1

Historia Natural

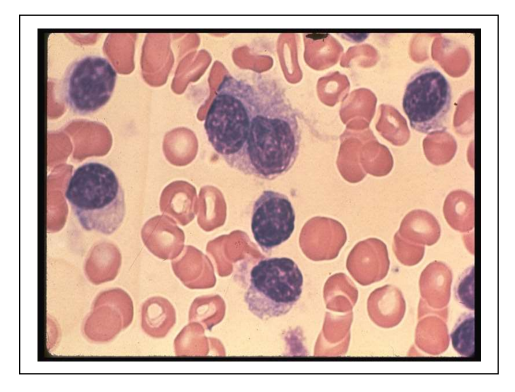
- Enfermedad indolente, supervivencia media: 11 años
- 1/3 muere por otras causas; 2/3 mueren de MW

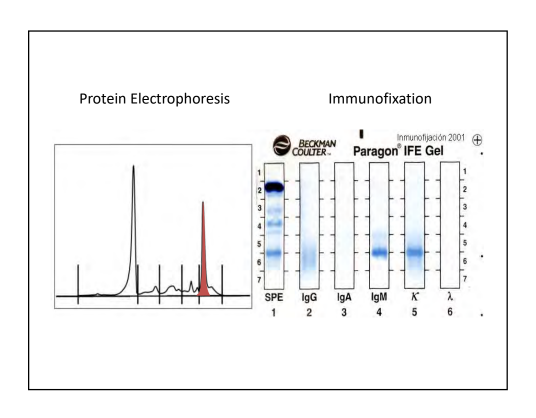
Tratamiento eficaz

Interés biológico

Definición de MW

- Trastorno linfoproliferativo raro caracterizado primariamente por infiltración de médula ósea y presencia de componente monoclonal IgM.
- El trastorno patológico subyacente en la MW es el linfoma linfoplasmocítico tal y como lo define las clasificaciones OMS [(WHO) World Health Organization] y REAL (Revised European-American Lymphoma).





Clasificación de Gammapatías IgM

	Proteína monoclonal IgM¹	Infiltración medular ²	Síntomas atribuibles a la lgM	Síntomas atribuibles a infiltración³
MW Sintomática	+	+	+ (4)	+ (4)
MW Asintomática	+	+	-	-
Trastorno tipo IgM ⁵	+	-	+	-
GMSI IgM	+	_	_	_

1) No se precisa un umbral de concentración de IgM para distinguir GMSI de MW, aunque la concentración de IgM raramente supera los 3 g/dL na GMSI, 2) Si un paciente tiene infiltración medular inequivoca por linfoma linfoplasmocítico, tiene MW; si no hay evidencias, se considerará GMSI. Sin embargo, hay pacientes con evidencias equivocas de infiltración de MO: ej, detección de células B clonales por citometría de flujo o PCR sin evidencias morfológicas de infiltración; o infiltrados equivocos de MO sin estudios confirmationos de cionalidad. En tanto no haya más datos, estos pacientes se calastifican como GMSI 3) Los sintomas atribuibles a infiltración tumoral son constitucionales, citopenias y organomegalias. 4) Se requiere la presencia de uno o ambos grupos de sintomas. (5) Población de pacientes que tienen sintomas atribuibles a la proteína monocional IgM, pero no tienen evidencias de células tumorales. Ej: crioglobulinemia sintomática, amiloidosis o fenómenos autoimunes tales como isquemia periférica y crioaglutininemia. Estos pacientes son un grupo clinicamente distinto y se propone el término "trastomos relacionados con IgM"

Owen et al. Sem Hematol 2003 30·110-115

Owen et al, Sem Hematol 2003, 30:110-115

BIOPSIA ÓSEA OBLIGATORIA

Essential evaluation of patients with Waldenström Macroglobulinaemia (WM)

Evaluation

- History and physical examination
- Include familial history for WM and other B-cell lymphoproliferative disorders
- Include funduscopic examination
- · Review of systems

Laboratory studies

- · Complete blood count
- · Complete metabolic panel
- Serum immunoglobulin levels (IgA, IgG, IgM)
- Serum and urine electrophoresis with immunofixation
- Serum beta-2-microglobulin level

If clinically indicated

- Cryoglobulins
- · Cold agglutinin titre
- · Serum viscosity
- · Screening for von Willebrand disease
- 24-h urine protein quantification

Bone marrow aspiration and biopsy Immunohistochemistry

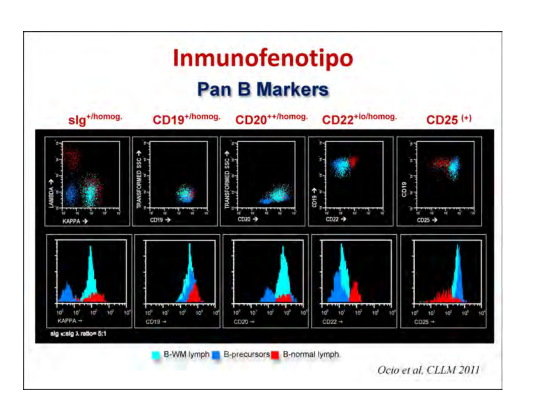
- · Flow cytometry
- Testing for MYD88 L265P gene mutation

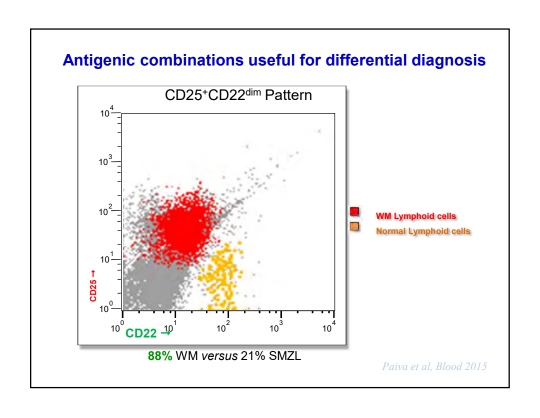
Computerized tomography scans of the chest, abdomen and pelvis with intravenous contrast

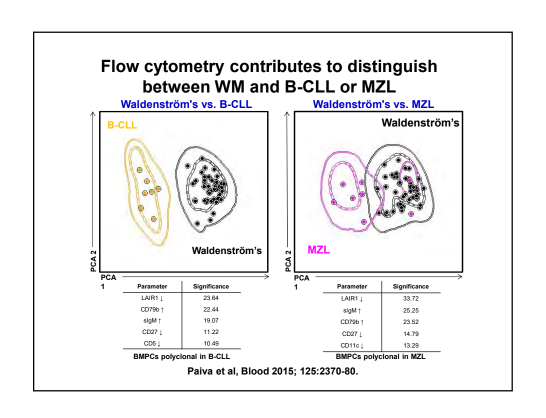
In patients being considered for therapy

Castillo JJ, Garcia-Sanz R, E Hatjiharissi, et al. Br J Haematol 2016, 175, 77–86

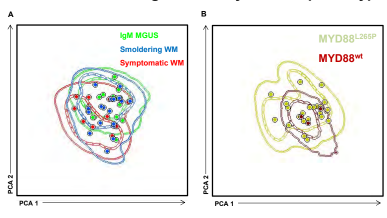
Son útiles el inmunofenotipo y las alteraciones citogenéticas y/o moleculares?







Waldenström's Macroglobulinemia MYD88 L^{265P} and MYD88 WT are indistinguishable by immunophenotyping



Paiva et al, Blood 2015; 125:2370-80.

Cytogenetic Abnormalities in WM

	Schop 2002	Ocio 2006	Fonseca 2006	Chang 2004	Nguyen- Khac 2010 N=132
Deletion of 6q21	42%	33%	47%	-	22%
Deletion of 13q14	16%	3%	-	9%	13%
Deletion of 17p23*	15%	7%	-	9%	8%
IgH translocations	0%	13%	2%	14%	3%
Deletion of 11q22	-	-	-	-	8%
Trisomy 4	-	-	-	-	8%
Trisomy 12	-	-	-	-	3%
Complex Karyotype (25/79)	-	-	-	-	32%
Trisomy 18 * Poor prognosis	-	-	-	-	11%



434 Whole-Genome Sequencing Results From 30 Patients with Waldenstrom's Macroglobulinemia

Hunter Z, Xu L, Zhou Y, Yang G, Liu X, Cao Y, Hanzis C, Sheehy P, Manning R, Patterson CJ, Laramie JM, Skifter DA, Lincoln SE, Treon SP Bing Center for Waldenstroms macroglobulinemia, Dana-Farber Cancer Institute, Boston MA, Boston, MA

The NEW ENGLAND JOURNAL of MEDICINE
N ENGL J MED 367;9 NEJM. ORG AUGUST 30, 2012

ORIGINAL ARTICLE

MYD88 L265P Somatic Mutation in Waldenström's Macroglobulinemia

Whole genome sequencing in WM

- Paired malignant (WM LPC) and normal (CD19-depelted PBMC) cell genomes from 10 WM patients were sequenced
- Genome from BM LPC alone were sequenced for additional 20 WM patients
- Results were validated by Sanger sequencing and included a cohort of WM, IgM MGUS, MM patients and healthy donors

Zachary Hunter ASH 2011, 434a

Results

Tumor and normal genomes were both sequenced to an average of 66X (range 60-91X) coverage of mapped individual reads. The average gross mapped yield for these genomes was 186.89 (range 171.56-262.03 Gb).

Acquired copy number changes were common:

- Losses: chromosome 6q (13/30; 43%),
- Gains:
 - -chromosome 4 (7/30; 23%)
 - -6p (3/30; 10%)
- Large regions of CNLOH were observed in 9/30 (30%) of patients occurring in chromosomes 1, 2, 3, 5, 9, 11, 17, 21, and X.

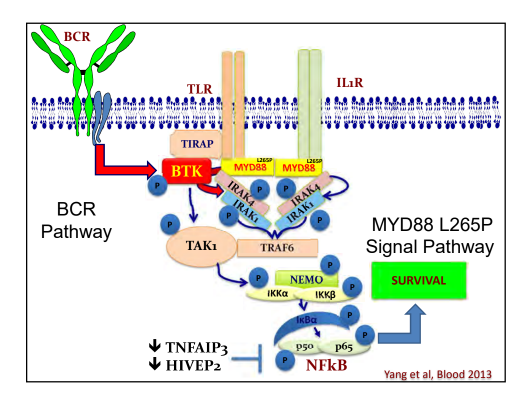
Results

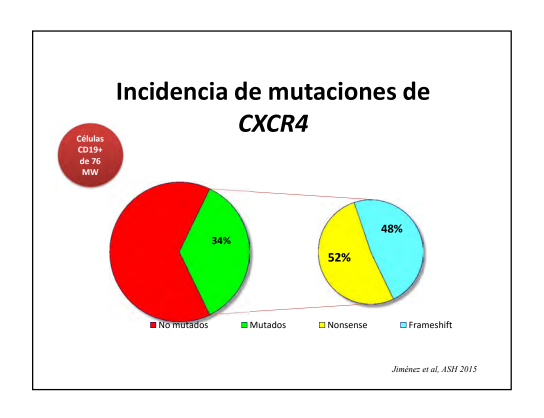
Involved Gene	Frequency
Myeloid differentiation primary response (MYD88) gene (38182641 in	26/30 (86.7%)
chromosome 3p22.2): 265 leucine → proline (L265P)	4/26 (15%) Homozygous, due to CNLOH
Transporter 2, ATP-binding cassette, sub-family B (TAP2) gene	7/30 (23%)
Chemokine (C-X-C motif) receptor 4 (CXCR4) gene	6/30 (20%)
Low density lipoprotein receptor-related protein 1B (LRP1B) gene	5/30 (17%)
Mesothelin (MSLN) gene	4/30 (13%)
AT rich interactive domain 1A (ARID1A)	3/30 (10%)
Histone cluster 1, H1e (HIST1H1E)	3/30 (10%)
Rap guanine nucleotide exchange factor 3 (RAPGEF3)	3/30 (10%)

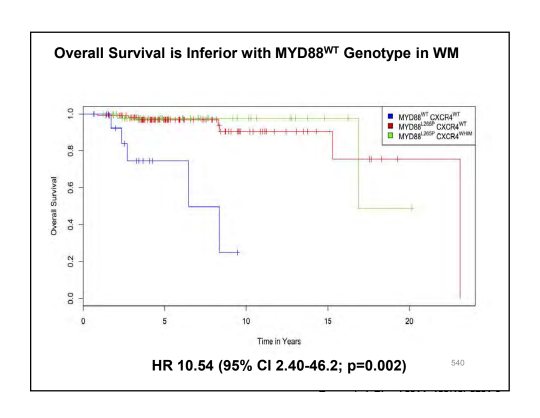
MYD88 L265P mutation in B-cell LPD

Entity	N	MYD88	L265P
Waldenström's Macroglobulinemia	117	101	(86%)
IgM MGUS	31	27	(87%)
Non-CG Diffuse Large Cell Lymphoma	48	9	(19%)
Marginal Zone Lymphomas	14	3	(21%)
B-CLL (18 with IgM M-component)	39	0	(0%)
Hairy cell Leukemia	36	0	(0%)
Lymphoplasmocytic lymphoma	9	0	(0%)
Multiple Myeloma (3 lgM)	24	0	(0%)
MGUS IgG/IgA	25	0	(0%)
Healthy volunteers	38	0	(0%)

Jiménez et al, Leukemia 2013







Existe algún índice pronóstico en MW?

Prognosis: Uni/Multivariate analyses: n=587

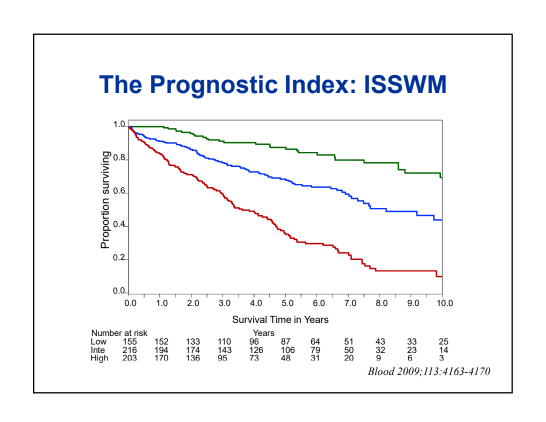
Characteristics	No of patients	Median survival	95%CI	p value
• Age				
<u>≤</u> 65	254	141	120-153	
> 65	333	56	49-63	< 0.0001
 B2M (mg/L) 				
≤3	251	122	103-141	
> 3	326	63	55-83	< 0.0001
 Hemoglobin (g/L) 				
≤ 11.5	381	123	110-179	
> 11.5	205	72	62-84	< 0.0001
Platelets (109/L)				
≤ 100	54	51	32-59	
> 100	531	90	83-116	< 0.0001
 Absolute neutrophil count (109/L) 				
≤ 1.5	53	46	27-74	
> 1.5	512	89	80-103	0.0018
 Serum monoclonal protein (g/L) 				
< 70	541	90	82-110	
> 70	43	49	37-62	0.0016
Serum albumin (g/L)				
< 35	197	79	55-89	
> 35	354	106	92-137	0.0012

Morel et al, Blood 2009;113:4163-4170

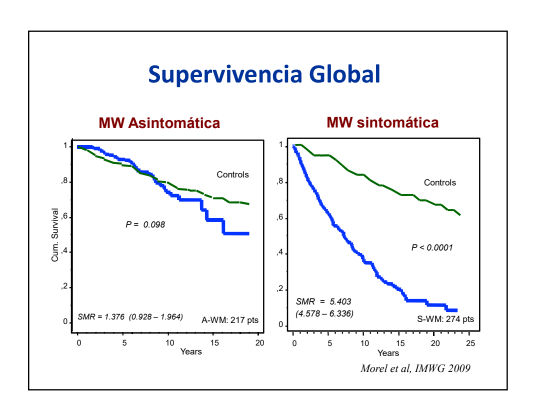
The Prognostic Index: ISSWM

Stratum	Score	Total	Failed	Median	0.95lcl	0.95ucl
Low	0 or 1 (except age)	155 (27%)	38	142.5	120.3	195.7
Intermediate	Age>65 or 2	216 (38%)	87	98.6	81.7	137.2
High	>2	203 (35%)	134	43.5	36.6	55.1

Morel et al, Blood 2009;113:4163-4170



Cómo lo trato?



Treatment Criteria: Symptomatic disease

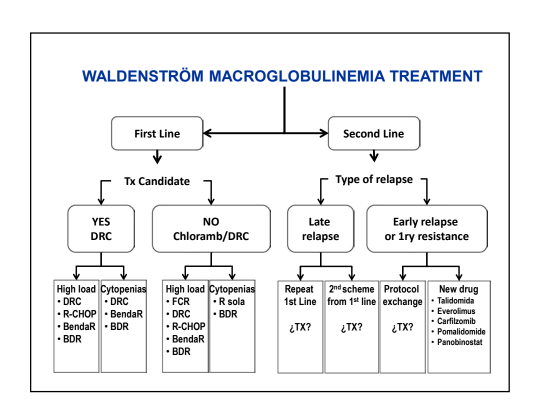
- 1. Recurrent fever, night sweats, weight loss, fatigue
- 2. Hyperviscosity
- Lympadenopathy which is either symptomatic or bulky (≥5cm in maximum diameter)
- 4. Symptomatic hepatomegaly and/or splenomegaly
- 5. Symptomatic organomegaly and/or organ or tissue infiltration
- 6. Peripheral neuropathy due to WM
- 7. Symptomatic cryoglobulinemia
- 8. Cold agglutinin anemia
- 9. Immune hemolytic anemia and/or thrombocytopenia
- 10. Nephropathy related to WM
- 11. Amyloidosis related to WM
- 12. Hemoglobin ≤10g/dL
- 13. Platelet count <100·109/L

Kyle et al. Semin.Oncol 2003; 30: 116-120

Therapeutic options

- ✓ Alkylators
 - Chlorambucil & prednisone
 - Chlorambucil continuous
 - Chlorambucil intermittent
 - COP
 - Melphalan & prednisona
- ✓ Polychemotherapy:
 - CHOP, M2, VAD
- ✓ Purine analogs
 - 2-Chloro-deoxi-adenosine
 - Fludarabine
 - 2-Deoxicoformicin

- ✓ Monoclonal Antibodies
 - Anti-CD20
 - Anti-CD52, Anti-CD22
- ✓ Proteasome inhibitors
 - Bortezomib
 - Cafilzomib
- ✓ IMiDs
- ✓ BTK inhibitors
- ✓ Transplant
 - Autologous
 - Alogeneic



Tratamiento de la macroglobulinemia de Waldenström RESUMEN						
Regimen	CR+PR	PFS	DR	os		
Alquilantes	40-70%	60 m	45 m	>5 años		
Fludarabina sola	50%	40 m	45 m	>5 años		
Rituximab solo	30%	46 m	72 m	>5 años		
Combo Análogos purinas	85-90%	45 m		>5 años		
R-CHOP	83%	62 m		>5 años		
CHOP	64%	18 m		>5 años		
CDR	83%	35 m		>5 años		
BendaR	96%	69 m		>5 años		
BDR	91%	45 m	66 m	>5 años		

Comparative Outcomes Following CP-R, CVP-R, and CHOP-R in Waldenström's Macroglobulinemia

Leukothea Ioakimidis, ¹ Christopher J. Patterson, ¹ Zachary R. Hunter, ¹ Jacob D. Soumerai, ¹ Robert J. Manning, ¹ Barry Turnbull, ² Patricia Sheehy, ¹ Steven P. Treon¹, ³

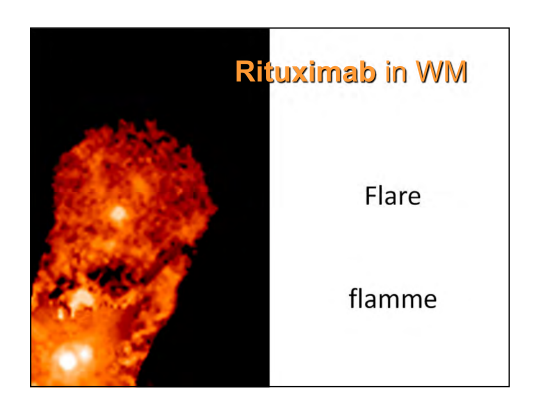
	N	ORR	CR	Comments
CHOP-R	23	96%	17%	Higher IgM (<i>P</i> = 0.015)
CVP-R	16	88%	12%	-
CP-R	19	95%	0%	↓ neutropenic fever and neuropathy

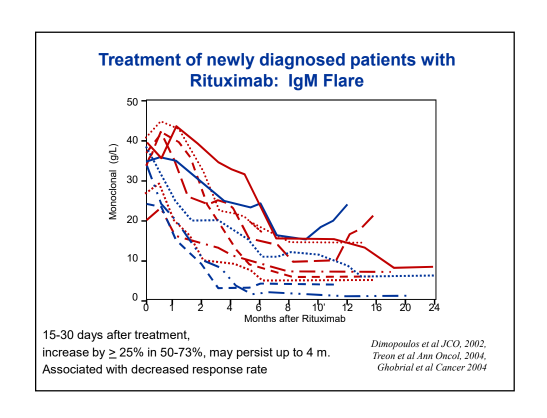
62 Clinical Lymphoma & Myeloma March 2009

Waldenström's Macroglobulinemia Treament in Previously untreated patients

Anti-CD20 and...

- 1. Alkylating agents
- 2. Nucleoside analogues
- 3. Immunomodulatory agents (IMiDs)
- 4. Bortezomib





DRC regimen

- Dexamethasone 20 mg IV day 1
- Rituximab 375 mg/m² IV day 1
- Cyclophosphamide 100 mg/m² PO BID days 1-5 (total dose 1000 mg/m²)

DRC courses are repeated every 21 days for 6 courses

Dimopoulos J Clin Oncol 2007; Kastritis Blood 2016

DRC regimen

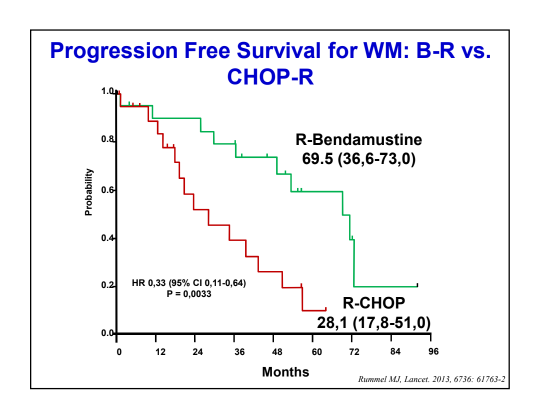
ORR = 83%

- N=72
- CR = 7%
- PR = 67%
-
- MR = 9%
- SD = 8%
- PD = 8%

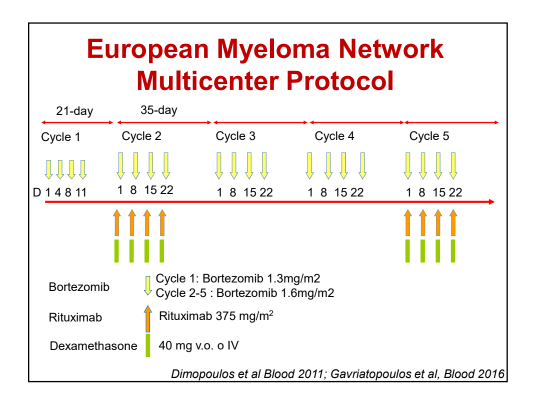
Median time to 50% IgM reduction was 4.1 months (range, 0.7-14)

IgM flare in 32%, >25% IgM increase in 11%

Dimopoulos J Clin Oncol 2007; Kastritis Blood 2016



	Autores	n	Ciclos	Asoc	RP	RAC
	Dimopoulos et al., Haematologica 05	10	6	No	60%	0%
sop	Chen et al., JCO 07	27	6	No	44%	0%
Iratados	Treon et al., CCR 07	27	6	No	48%	0%
G	Ghobrial et al, JCO 10	37	6	R	87%	5%
No tratados	Treon et al., JCO 2009*	23	6	RD	96%	22%
No IIa	Dimopoulos, Blood 2013*	59	6	RD	85%	10%



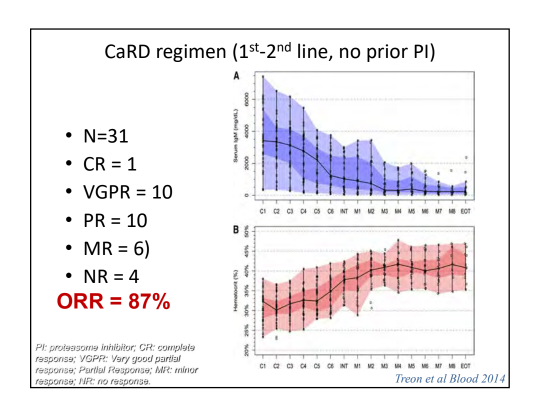
CaRD

- Induction
 - Carfilzomib IV,
 - 20 mg/m², infused over 20 minutes (cycle 1 only)
 - 36 mg/m² infused over 30 minutes (for cycles 2 and beyond)
 - Dexamethasone IV, 20 mg, given on days 1-2 & 8-9
 - Rituximab, 375 mg/m², days 2 and 9

Every 21 days for 6 cycles

- Maintenance (for stable disease or better), 8 weeks later:
 - Carfilzomib IV, 36 mg/m², D1 & D2
 - Dexamethasone IV, 20 mg, D1 & D2
 - Rituximab, 375 mg/m2, D2 only

Every 8 weeks for 8 cycles.



	CaRD, toxicitity							
	Car	D, LOX	icitity					
			-					
Table 2. Adverse events possibly, probably, or definitely associated with protocol therapy								
Toxicity type	Any grade	Grade 1	Grade 2	Grade 3	Grade 4			
Anemia	3 (9.7%)	0 (0%)	2 (6.5%)	1 (3.2%)	0 (0%)			
Arthralgia	3 (9.7%)	3 (9.7%)	0 (0%)	0 (0%)	0 (0%)			
Azotemia	3 (9.7%)	1 (3.2%)	2 (6.5%)	0 (0%)	0 (0%)			
Cardiomyopathy	1 (3.2%)	0 (0%)	0 (0%)	1 (3.2%)	0 (0%)			
Chest pain (non-cardiac)	1 (3.2%)	0 (0%)	1 (3.2%)	0 (0%)	0 (0%)			
Dyspepsia	1 (3.2%)	1 (3.2%)	0 (0%)	0 (0%)	0 (0%)			
Fatigue	2 (6.5%)	1 (3.2%)	1 (3.2%)	0 (0%)	0 (0%)			
Hyperglycemia	31 (100%)	7 (22.6%)	17 (54.8%)	7 (22.6%)	0 (0%)			
Hyperamylasemia	8 (25.8%)	7 (22.6%)	1 (3.2%)	0 (0%)	0 (0%)			
Hyperbilirubinemia	9 (29.0%)	7 (22.6%)	2 (6.5%)	0 (0%)	0 (0%)			
Hyperkalemia	1 (3.2%)	1 (3.2%)	0 (0%)	0 (0%)	0 (0%)			
Hyperlipasemia	17 (54.8%)	4 (12.9%)	8 (25.8%)	5 (16.1%)	0 (0%)			
Hypokalemia	1 (3.2%)	1 (3.2%)	0 (0%)	0 (0%)	0 (0%)			
Insomnia	2 (6.5%)	1 (3.2%)	1 (3.2%)	0 (0%)	0 (0%)			
Infusion reaction (rituximab)	7 (22.6%)	1 (3.2%)	6 (19.4%)	0 (0%)	0 (0%)			
Mucositis	2 (3.2%)	2 (3.2%)	0 (0%)	0 (0%)	0 (0%)			
Neutropenia	11 (34.8%)	7 (22.6%)	1 (3.2%)	2 (6.5%)	1 (3.2%			
Peripheral neuropathy	6 (19.4%)	5 (16.1%)	1 (3.2%)	0 (0%)	0 (0%)			
Rash	9 (29.0%)	6 (19.4%)	3 (9.7%)	0 (0%)	0 (0%)			

2956 Ixazomib, Dexamethasone and Rituximab in previously untreated patients with Waldenstrom Macroglobulinemia (Castillo JJ)

Phase II

Induction: six 4-week cycles

Ixazomib 4 mg 1, 8, 15 Dexamethasone 20 mg 1,8,15 Rituximab 375 mg/m2 IV 1

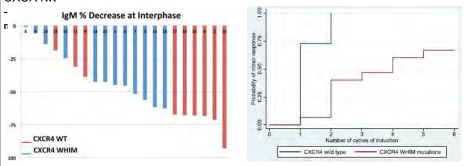
Maintenance: Six 8-week cycles

Characteristic	Patients (N=26)	
Age at WM diagnosis – yr	63 (46-81)	
Age of treatment initiation – yr	65 (46-82)	
Hemoglobin – g/dL	10.2 (6.9-13.2)	
Serum IgM – mg/dL	5,068 (653- 7,650)	
Bone marrow involvement - %	55 (5-95)	
Lymphadenopathy – no. (%)	46 (12)	
Splenomegaly	12(3)	
MYD88 L265P	100 (26)	
CXCR4 WHIM	58 (15)	
Nonsense	67 (10)	
Frameshift	33 (5)	
Criteria for treatment initiation - no.		
(%)	13 (48.1)	
Anemia	1 (3.8)	
Symptomatic splenomegaly	7 (27)	
Hyperviscosity	4 (15.4)	
Peripheral neuropathy	5 (19.2)	
Constitutional symptoms	1 (3.8)	
IgM >6,000 mg/dL Pancytopenia	1 (3.8)	

2956 Ixazomib, Dexamethasone and Rituximab in previously untreated patients with Waldenstrom Macroglobulinemia (Castillo JJ)

Results: (of the 16 patients completed induction)

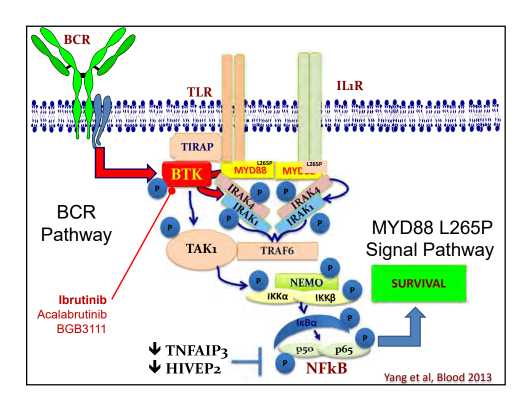
- -Median Time To Response : 8 weeks
- -ORR: 80% (VGPR 5%, PR 45%, MR 30%)
- -The median time to Minor Response in CXCR4mut was 4 cycles vs 1 cycle en CXCR4wt
- -Major response (VGPR+PR) were observed in 40% CXCR4mut patients vs 55% CXCR4wt



Conclusion: IDR active, well tolerated, no PN. CXCR4 mutation affect time /dept of response

Waldenström's Macroglobulinemia Treatment in Previously treated patients

- "Late" Relapses after initial response (>12 months)
 - Same prior line or another 1st line protocol
- "Early" relapses after initial response, and primary or secondary relpases:
 - Switch for another first line
 - "Conventional" Lymphoma strategy:
 - Polychemotherapy (plus R): CHOP, CAP, VBCMP y VAD
 - Trasplant: Auto or Alo
 - New therapies ("experimental"):
 - Bortezomib, IMiDs, Alemtuzumab, Bendamustine...
 - Everolimus
 - · Carfilzomib
 - Pomalidomide
 - Panobinostat



Ibrutinib in Previously Treated Waldenstrom's Macroglobulinemia

Steven P. Treon, Christina Tripsas, Kirsten, Diane, Guarav Varma, Rebecca Green, Kimon Argyropoulos, Guang Yang, Yang Cao, Lian Xu, Christopher J. Patterson, Scott Rodig, James L. Zehnder, Jon C. Aster, Nancy Lee Harris, Sandra Kanan, Irene Ghobrial, Jorge Castillo, Jacob Laubach, Zachary R. Hunter, M. Lia Palomba, and Ranjana Advani.

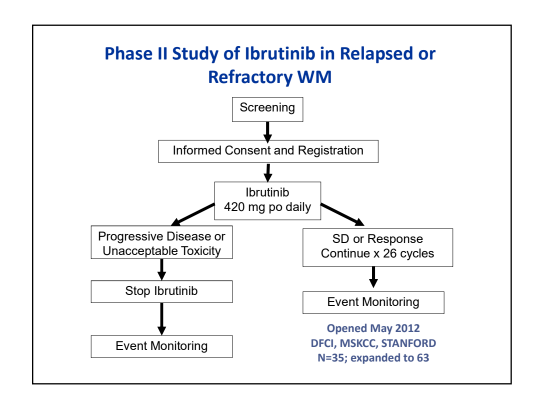
Dana-Farber Cancer Institute Memorial Sloan-Kettering Cancer Center Stanford University Medical Center





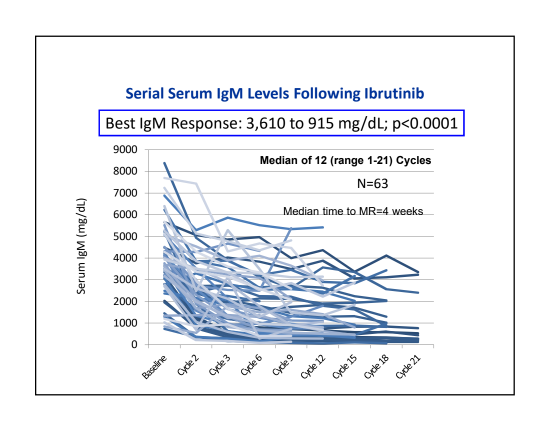


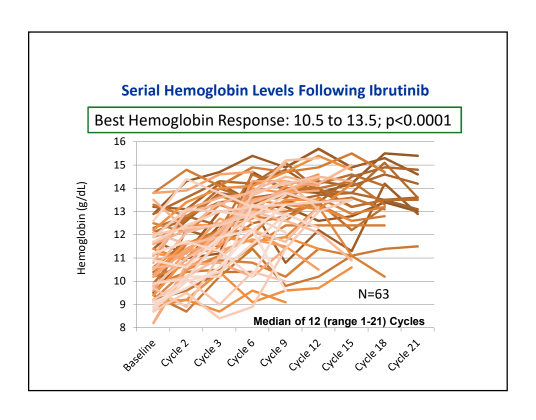
Treon et al NEJM, 2015



Baseline Characteristics for Study Participants (n=63)

Characteristic	Median	Range	
Age (yrs)	63	44-86	
Male/Female	48/15	N/A	
Prior therapies	2	1-8	
Hemoglobin (mg/dL)	10.5	8.2-13.8	
Platelet (k/uL)	214	24-459	
Serum IgM (mg/dL)	3,610	735-8,390	
B ₂ M (mg/dL)	3.9	1.3-14.2	
BM Involvement (%)	70	3-95	
Adenopathy >1.5 cm	37 (58.7%)	N/A	
Splenomegaly >15 cm	7 (11.1%)	N/A	



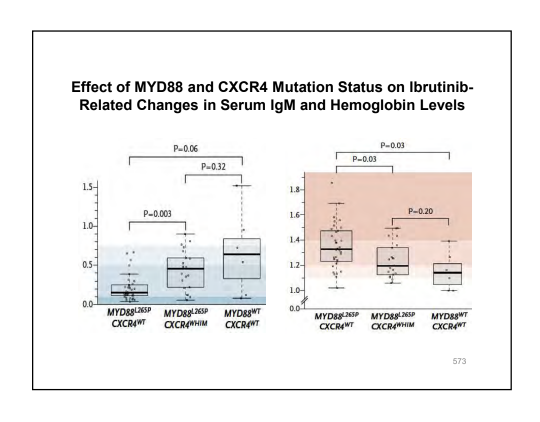


Best Clinical Responses to Ibrutinib

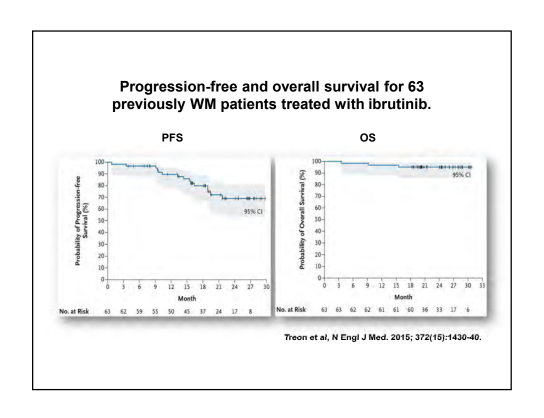
Median of 12 (range 1-21) Cycles

	(N=63)	(%)
VGPR	9	14
PR	34	54
MR	12	19
SD	7	11
Non-Responder	1	2

ORR: 87% Major RR (≥PR): 68%



MYD88 (•	brutinib are i on-L265P) and		
	MYD88 ^{MUT} CXCR4 ^{WT}	MYD88 ^{MUT} CXCR4 ^{WHIM}	MYD88 ^{WT} CXCR4 ^{WT}	p-value
N=	36	21	5	
Overall RR	100%	85.7%	60%	<0.01
Major RR	91.7%	61.9%	0%	<0.01

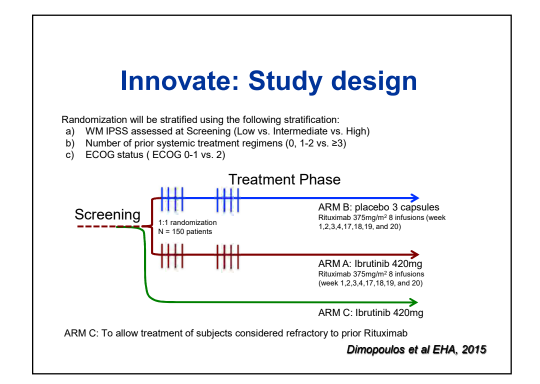


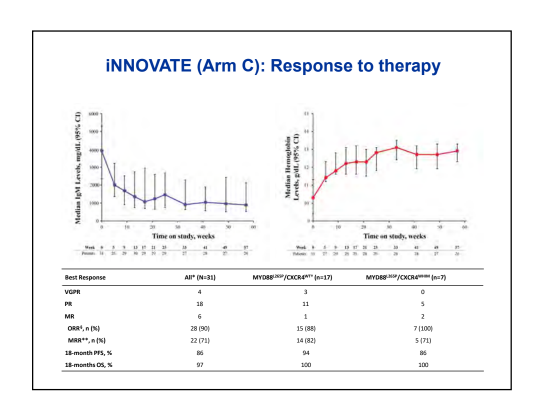
		a march	se Event	
	Grade 2	Grade 3	Grade 4	Total Grade 2-
Blood and lymphatic system disc			13.000	1
Anemia	9 (14.3%)	3 (4.8%)	0 (0%)	12 (19.0%)
Neutropenia	5 (7.9%)	6 (9.5%)	5 (7.9%)	16 (25,4%)
Thrombocytopenia * Cardiac disorders	2 (3.2%)	5 (7.9%)	2 (3.2%)	9 (14.3%)
Atnal fibrillation	2 (3.2%)	1 (1.0%)	0 (0%)	3 (4.8%)
Sinus tachycardia	1 (1.0%)	0 (0%)	0 (0%)	1 (1.0%)
Gastrointestinal disorders	131333	3.40 10.2		1
Dianhea	3 (4.8%)	0 (0%)	0 (0%)	3 (4.8%)
Gastroesophageal reflux disease	1.(1.0%)	D (0%)	0 (0%)	1 (3.0%)
Mucositis oral	2 (3.2%)	0 (0%)	0 (0%)	2 (3.2%)
Infections and infestations				
Febrile neutropenia	0.(0%)	O (0%)	1 (1.6%)	1 (1.0%)
Endocarditis infective	0 (0%)	1 (1.0%)	0 (0%)	1 (1.6%)
Lung infection	4 (6.3%)	0 (0%)	0 (0%)	4 (6.3%)
Sinusitis	1 (1.6%)	D (0%)	D (0%)	1 (1.6%)
Skin infection Urinary tract infection	3 (4.8%)	1 (7.6%)	D (0%)	4 (6.3%)
Injury, poisoning and procedural		D (D%)	D (Q%)	1 (5/6%)
Postoperative hemorrhage	1 (1.0%)	10 (0%)	0 (0%)	1 (1.0%)
Metabolism and nutrition disords		2007	0.00.47	117.000
Dehydration	2 (3,2%)	0 (0%)	0.(0%)	2 (3.2%)
Musculoskeletal and connective				
Arthraigia	1 (1:6%)	D (0%)	0 (0%)	1 (1.6%)
Nervous system disorders				-
Presyncope Syncope	1 (1.6%)	D (0%)	0 (0%)	1 (1.6%)
Respiratory, thoracic and medias	0 (0%)	1 (1.8%)	0 (0%)	1 (1.0%)
Epistaxis	2 (3.2%)	0 (0%)	-0 (0%)	2 (3.2%)
Cough	1 (1.6%)	D (0%)	D (0%)	1 (1.6%)
Skin and subcutaneous tissue di				
Pruritus	1 (1:0%)	10 (0%)	0 (0%)	1 (1.0%)
Folijouittis	1 (1.6%)	D (0%)	D (0%)	2 (1.6%)
Rash	7 (1.6%)	D (0%)	0 (0%)	1 (1.0%)
Skin Peeling	1 (1/6%)	0 (0%)	0 (0%)	1 (1/6%)
Vascular disorders		111 860	W Second	T. A SA MARY
Hematoma Hypertension	0 (0%)	1.(1.6%)	0 (0%)	1 (1.6%)
Hypotension	2 (3.2%)	D (0%)	0 (0%)	1 (1.6%)
rypotension	1.[1.0/36]	W 20176-L	0 (0%)	1.(1.034)

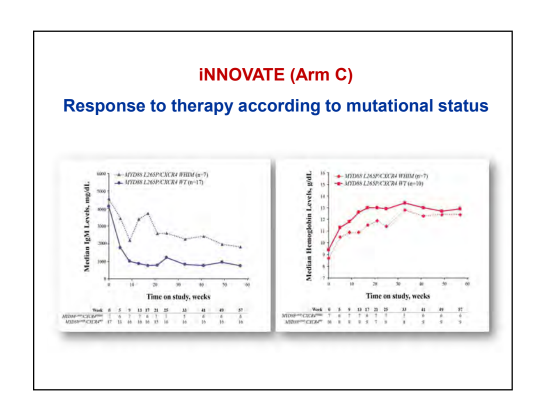
Single-Agent Ibrutinib in Rituximab-Refractory Patients with Waldenström's Macroglobulinemia: Results From a Multicenter, Open-Label Phase 3 Substudy (iNNOVATE™)

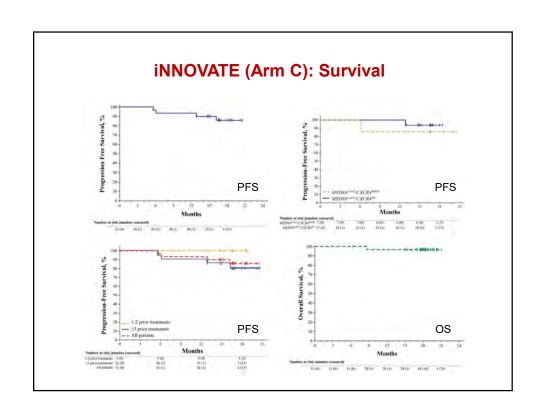
Dimopoulos, MA, Trotman J, Tedeschi A, Matous JV, Macdonald D, Tam C, Tournilhac O, Ma S, Oriol A, Heffner LT, Shustik C, García-Sanz R, Cornell RF, Fernández de Larrea C, Castillo JJ, Granell M, Kyrtsonis MC, Leblond V, Symeonidis A, Kastritis E, Singh P, Li J, Graef T, Bilotti E, Treon S, Buske C, on behalf of the iNNOVATETM Study Group and the European Consortium for Waldenström's Macroglobulinemia (ECWM).

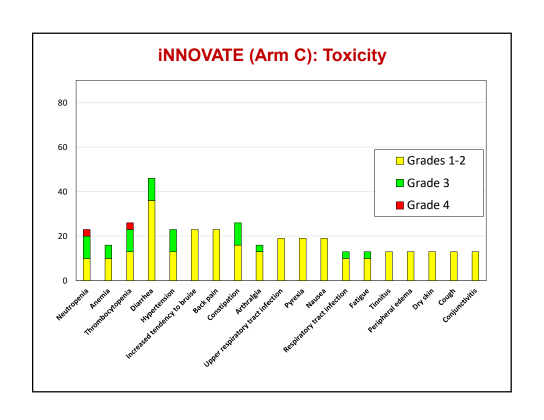
Dimopoulos Lancet Oncol. 2017 Feb;18(2):241-250











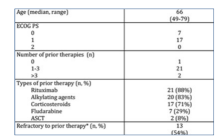
ASH2016: #1216 High Major response Rate, Including VGPR, in Patients with Waldenstrom Macroglobulinemia (WM) treated with the Highly Specific BTK inhibitor Bgb-3111:Expansion Phase Results from ongoing Phase I Study (Tam CS)

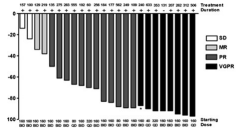
Introduction: BGB-3111 irreversible BTK inhibitor with greater selectivity and superior bioavailability

Phase I (6 ptes) 40,80,160,320 mg and Phase 2 (25 ptes) 160, 320 mg; **24 evaluable for response**

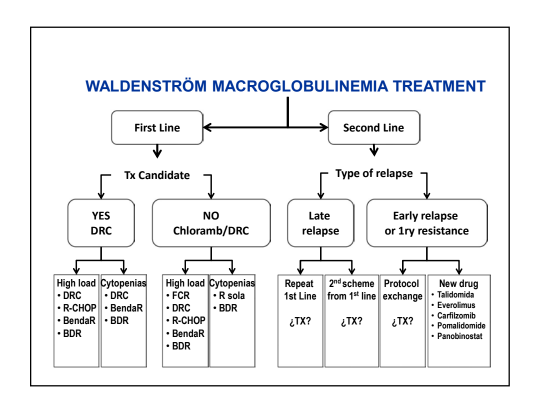
Major Response 83% (VGPR 33%, PR 50%)

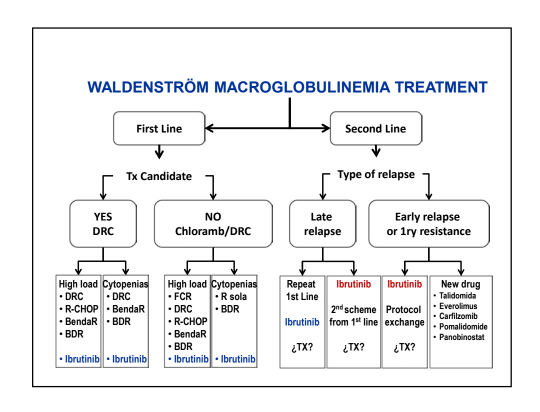
AE's (1/2). Respiratory infection 25%, diarrhea 25% and nausea 21%

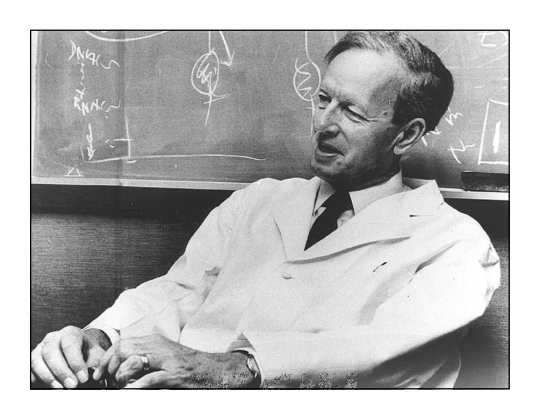




Conclusions: BGB-3111 is well tolerated and highly active in WM. Te depth and quality of response, warrant a randomized comparison against Ibrutinib









TREATMENT OF DISEASE COMPLICATIONS

Jesús G. Berdeja, M.D. Director of Multiple Myeloma Research Sarah Cannon Research Institute Tennessee Oncology Nashville, TN, USA

disclosures

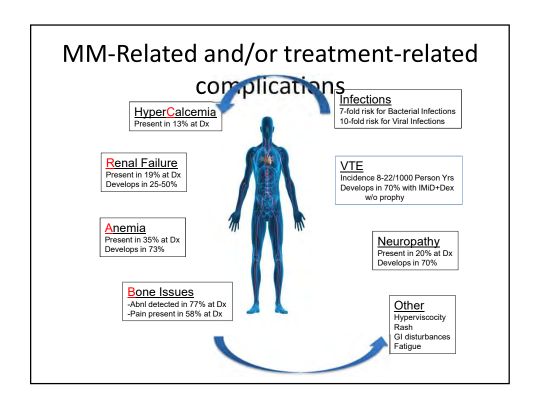
- Research funding from the following:
 - Abbvie, Amgen, Bluebird, BMS, Celgene, Janssen, Novartis, Takeda, Teva

AIMS

- Identify complications due to MM and its treatments
- Review standard recommendations for prevention of complications
- Recommended management of complications related to MM itself
- Review standard recommendations for management of treatment-related complications

Case: Presentation

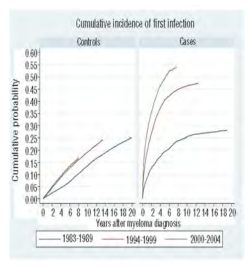
- A 69-yr-old man presents with low back pain, fatigue
- Physical examination: T 36.4°, BP 186/79 mm Hg, P 75/min, R 20/min
- · Laboratory assessments
 - Hb 9.6 g/dL
 - Serum creatinine 2.35 mg/dL
 - Calcium 11.9 mg/dL
 - IgG 3800 mg/dL, IgA 43 mg/dL, IgM 9 mg/dL
 - SPEP 3gm/dL abnormal protein, IFE IgGk
 - SFKLC 200 mg/dL
 - $\quad \beta_2\text{-microglobulin 6.9 mg/L}$
 - Albumin 3.2 g/dL
 - LDH 200 nl
- Skeletal survey: small lytic lesions spine and pelvis, compression fracture at T12
- Bone marrow biopsy: 70% k-restricted plasma cells



Immunosuppression in mm

- MM leads to multifactorial humoral and cellular immune dysfunction
- Anti-myeloma therapy is immunosuppressive
 - PIs can suppress T cell function
 - Grade 3 /4 infections in 35% of newly dx pts reported in VRD arm in SWOG S0777 trial
 - Lenalidomide and pomalidomide can cause neutropenia in >25% patients
 - Grade 3 /4 infections in 29% newly dx pts on len/dex reported by FIRST trial
 - Dexamethasone is pan-immmunosuppressive
- Ptts://differentiages/2021 Principles of the Principles of th





Cecilie Blimark et al. Haematologica 2015;100:107-113

Take home points:

- -7-fold risk of bacterial infection
- -10-fold risk of viral infection
- -Infectious risk greatest during 1st yr dx
- -Infections account for 20% early mortality highest risk first 2-4 mos of treatment
- -Bacterial infections predominate during the first weeks of initial therapy
- -Viral infections, often due to reactivation and most common during PI/Dex treatment

recommendations

- Infection prophylaxis
 - Pts should remain up to date on appropriate vaccinations
 - Yearly influenza recommended
 - Pneumovax recommended though unclear benefit but if done should be done at time of best disease control
 - Live vaccines should be avoide
 - VZV prophylaxis when receiving PIs, monoclonal abs, during ASCT recovery period is recommended with acyclovir or similar agents
 - Antibacterial prophylaxis with mixed results
 - Not routinely recommended
 - Consider in high risk patients at initial dx 2-3 mos
 - Consider with lenalidomide/pomalidomide therapy during 1st 3 mos
 - Use of IVIG is controversial
 - Consider in pts with hypogammaglobunemia and recurrent infections
 - Consider use of G-CSF in treatment induced neutropenia or neutropenic fever

Terpos et al: Haematologica 2015. Anderson et al: J Natl Compr Cancer Netw 2016. Chanan-Khan et al: JCO 2008. Lonial et al: NEJM 2015. Oken et al: Am J Med 1996. Vesole et al: Leukemia 2012. Chapel et al: Lancet 1994. Musto et al: BJH 1995. Aapro et al: EurJCan 2011.

Bone disease: scope of problem

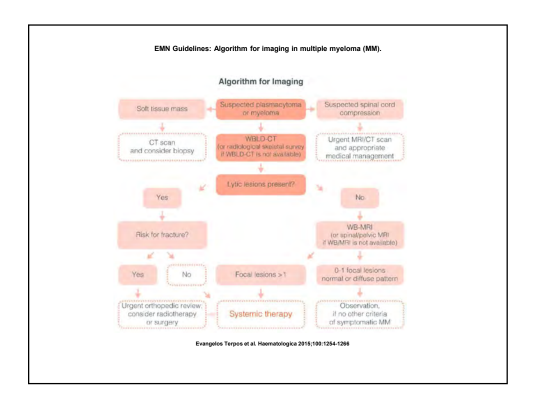
- Osteolytic bone disease is present in up to 80% of pts at dx
- Increased osteoclast activity leads to skeletalrelated events (SREs)
 - Vertebral CF
 - Other pathologic fractures
 - Spinal Cord Compression

Terpos et al: Ann Oncol 2005. Kyle et al: Mayo Clin Proc 2003.

imaging

- Skeletal Survey (WBXR)
 - Current standard technique for detection of lytic lesions, widely available
- WBLD-CT
 - More sensitive and accurate than WBXR and easy to perform where available
 - Used more and more in parts of the world as a replacement for WBXR
- WB-MRI
 - now indicated as part of the definition of active MM
 - >1 focal lesion by MRI is a MM-defining
 - depicts bone marrow involvement
- PET/CT May be substituted for MRI as above
 - May correlate better with response to therapy

Dimopoulos et al. 3 eurlogia 2092 frightinet N/TRTD appearate 2614 Welker et al.: JCO 2007. Waheed et al.: Haematologica 2013. Hillengass et al. JCO 2010. Kastritis et al.: Leukemia 2014. Zamaggin et al.: Blood 2011. Spinnato et al.: EurJRadiol 2012. Derlin et al.: Eur Radiol 2013.



Bisphosphonate use

- Pamidronate and zoledronic acid (ZA)have been found to reduce SREs in Phase III studies
- IV ZA >>> PO Clodranate
 - Reduced SREs
 - Improved OS by 10 mos in MM pts with lytic lesions
 - Effects continued beyond 2 yrs but unknow if all depths of remission benefitted equally
- No study has shown superiority of ZA over pamidronate but a meta-analysis showed a survival advantage to ZA over place bood 2012. Mhaskar et al.: CDSR 2012.

Side effects of bisphosphanates

- Acute phase reactions
- Hypocalcemia
- Hypophosphatemia
- Renal impairment
- Osteonecrosis of the jaw (ONJ)

Terpos et al: JCO 2013. Berenson et al: JClinPharmacol 1997; Dimopoulos et al: Haematologica 2006.

EMN/IMWG Recommendations for bone health

- ZA 4mg over 15-30 min or pam 90mg over 2-4 h IV q3-4wks
 - All MM pts with CrCl >30 mL/min and osteolytic lesions
 - Continue ZA past 2 yrs though unclear if benefit continues for pts in ≥ VGPR, pam unclear benefit past 2 yrs
- Calcium po 600mg/d; vitamin D2 400 IU/d
 - 60% of MM pts are Vitamin D deficient, yearly levels recommended
- Adjust ZA for renal fx, hold until crcl within 10% baseline
 - hold for crcl < 30. HD ok if no expectation of recovery
- ONJ more common with ZA, after invasive dental procedures, and associated with prolonged use of ZA
 - Try to have dental clearance and extractions/traumatic procedure prior to initiating bisphosphonates
 - IMWF recommend holding for 90 days pre and post invasive dental procedure

Terpos et al: Haematologica 2015. Terpos et al JCO 2013.

Bone pain treatment

- Radiotherapy for solitary plasmacytoma, symptomatic SC compression, severe uncontrolled pain or prevention of pathologic fx
 - 3000 cGy in 10-15 fx usually sufficient
 - Can lead to delays of systemic therapy
 - Can adversely affect stem cell mobilization
- · Vertebroplasty vs Kyphoplasty
- Surgery rare
 - For pathologic fractures of long bones
 - Restore spinal stability

Niesvizky et al: J Natl Compr Canc Netw 2010. Christoulas et al: Expert Rev Hematol 2009.

Drake MT. Oncology (Williston Park) 2009. Terpos et al: JCO 2013. Webb et al: Br J Pharmacol 2014.

VERTEBROPLASTY VS KYPHOPLASTY





- Height loss
 Upright posture becomes impossible
 Pulmonary volume loss due to
- anterior wedging of the spine d. 12 rib rests on the iliac crest e. Narrowed gab between ribs and ilium
- f. Protruding abdomen
- g. Distension, constipation, early satiety, eructation

Berenson et al: Lancet Oncol 2011. Bhargava et al: JCO 2009

Renal impairment – Defined as GFR < 60mL/min

- Present in 19% of pts at dx
- Develops in 25-50% at some point in disease course
- Pathophysiology is complex
 - Principal renal mechanism is cast nephropathy
 - Direct consequence of high serum concentration of SFLC
 - Direct deposition of light chains as in LCDD or amyloidosis
- · Other offending conditions
 - Hypercalcemia
 - Dehydration
 - Nephrotoxic medications

Kebler et al: Eur J Haematol 2009. Dimopoulos et al: Leukemia 2008. Hutchison et al: Nat Rev Nephrol 2011.

Management

- MM patient with RI should be considered an emergency
- Interventions should include
 - Hydration +/- urine alkalization
 - treatment of hypercalcemia
 - prompt anti-myeloma therapy
 - Use of plasma exchange, standard dialysis, high cut off dialyzers are reasonable when available but have not been proven to definitive impact course and thus not

Burndite COMMON ended nasestiandary dealtathistim annetti et al: Am J Hematol 2015.

Antimyeloma therapy in pts with renal impairment - frontline

- Bortezomib-based combinations have been shown to be most effective in reversing renal insufficiency and are the treatment of choice.
 - BTZ-based therapy may even overcome negative prognostic effect of RI (HOVON-65 trial)
- Bortezomib/Dex + Thalidomide (VTD) or cyclophosphamide (CyBorD) or doxorubicin (PAD) do not require dose-adjustments and should be 1st line
- In elderly patients also consider bortezomib, melphalan, prednisone (VMP)
- Lenalidomide can be used as long as the recommended adjustments for 2010 pimpowles state of the special state of the special state of the special special state of the special speci

Antimyeloma therapy in pts with renal impairment – transplant

- High dose melphalan and ASCT can be performed safely in patient with RI, including on dialysis
- Traditionally melphalan dose has been reduced to 140mg/m2 in patients with RI
- Recent analysis by the CIBMTR reported improved outcomes and no worse toxicity in patients with mild-moderate RI (CcCl > 30mL/min) when treated with melphalan

Badro 200 (1942) 12 tal: EJH 2005. Lee et al: BMT 2004. Raab et al: Haematologica 2006. Sweiss et al: BMT 2016. Mahindra et al: ASH

Other Antimyeloma therapies in pts with renal impairment

- Pomalidomide
 - No need to adjust, at least with CrCl >30mL/min
- Elotuzumab
 - No need to adjust even on HD
- Carfilzomib
 - No need to adjust
- Daratumumab
 - No data but likely safe?
- Bendamustine
 - no need to adjust
- Ixazomib

weiselet needs adjustment only for pits, with Cr Clis 30 ml/min or on HD16.

Case 2: Presentation

- 57-yr-old man diagnosed with MM
- Was started on
 - Bortezomib
 - 1.3 mg/m² IV on Days 1, 4, 8, 11
 - Thalidomide
 - 100 mg PO on Days 1-21
 - Dexamethasone
 - 40 mg PO on day of and day following btz
- After cycle 2, M-spike had decreased to 0.3 g/dL, but noted numbness, burning, stabbing sensation feet and legs

PERIPHERAL NEUROPATHY: RISK FACTORS AND GENERAL CONSIDERATIONS

COMMON PROBLEM AT BASELINE DUE TO CO-MORBIDITIES AND MM AND EVENTUALLY FROM

General Considerations

- Endocrine disorders
 - Hypothyroidism
 - Diabetes
- Nutritional disease
- Connective tissue disease
- Vascular disease
- Medications
- Herpes zoster
- Most common symptoms
 - Sensory deficits
 - Neuropathic pain

Disease- and Treatment-Related Factors

- Hyperviscosity syndrome
- Hypergammaglobulinemia
- Incidence of peripheral neuropathy at diagnosis: 20%
- Up to 75% of pts on therapy
- Incidence of grade 3/4 CIPN with novel agents
 - Bortezomib: 6% to 22%
 - → with wklv vs twice-weekly dosing
 - → with SC administration
 - Thalidomide: 3% to 23%
 - \(\bullet\) with higher doses and prolonged therapy, often permanent
 - Carfilzomib: < 2%

Gleason C, et al. J Natl Compr Cancer Netw. 2009;7:971-979. Palumbo A, et al. J Clin Oncol. 2014;32:587-600. Kurtin S, et al. J Adv Pract Oncol. 2013;4:307-321. Siegel D, et al. Haematologica. 2013;98:1753-1761. Richardson et al: Leukemia 2012. Delforge et al: Lancet Oncol 2010.

Thalidomide VS Bortezomib

PN and associated symptoms

- Thalidomide
 - Affects larger myelinated axons
 - Cumulative
 - Dose-dependent
 - Mostly irreversible
 - Autonomic dysfunction
 - Dizziness
 - Orthostasis
 - Tremor
 - Reversible with dose adjustment/discontinuation

- Bortezomib
 - Affects small myelinated and unmyelinated fibers
 - Hyperesthesia
 - Neuralgia
 - Altered temperature sensation
 - Reversible if intervene early
 - Autonomic dysfunction

Richardson et al: Leukemia 2012. Delforge et al: Lancet Oncol 2010. Delforge et al: Blood 2017

PN management

- Careful attention with pts on therapy
 - Prompt dose reductions for thalidomide
 - Bortezomib SQ dosing, weekly dosing, dose reduction
 - · Randomized trial IV vs SQ dosing
 - all grade PN decreased from 53% to 38%
 - grade 3 and 4 PN decreased from 16% to 6%.
- Treat neuropathic pain (ref)
 - Analgesics
 - Calcium channel blockers (e.g. gabapentin, pregabalin)
 - Serotonin-norepinephrine reuptake inhibitors
 - Tricyclic antidepressants (e.g. amitriptyline)
 - Acetyl-L-carnatin and alpha lipoic acid have shown activity
 Rithrethethtosindiaced PN toutorico prospective 2 analyses J Neurooncol 2015

Venous Thromboembolism (Vte)

- MM patients at higher risk due to several factors specific to disease and treatments
- Incidence VTE is 8-22/1000 person years
- Highest risk during first 4 months of treatment
- Incidence increases by up to 70% with IMiDs+Dex in absence of anticoagulation
- A randomized study with lenalidomide-based therapy established benefit of thromboprophylaxis with both ASA and enoxaparin
 - The incidence of VTE was
 - 2.3% in patients on aspirin 100mg/day
 - 1e. 2% rimi et Senon i emicka es 27 în 240 migs da yeukemia 2011. Dimopoulos et al: Leukemia 2014. Palumbo et al: Leukemia 2008 Larocca et al: Blood 2012.

Imwg risk factor assessment and					
prophylaxis recommendations ia. 2008,22:414-423.					
Treatment-related	Patient-specific	Myeloma-specific			
IMiDs	Age	Active/uncontrolled Disease			
High-dose Dexamethasone	Previous VTE	Hyperviscocity			
Erythropoietin	Infection				
Anthracyclines	Surgical Procedure				
Multiagent chemotherapy	Cardiovascular Co-morbs				
	Immobilization				
	Inherited thrombophilia				
	Central venous catheter				
	Recommendations for thromboprophylaxis				

Therapy

ASA

ASA

LMWH or warfarin

LMWH or warfarin

Risk Assessment for VTEs in Pts Receiving IMiD-based Therapy

- VTE prophylaxis for individual risk factors (eg, age or obesity) or myeloma-related risk factors (eg, immobilization or hyperviscosity)
 - If ≤ 1 risk factor present, aspirin 81-325 mg/day

Number of risk factors

≥ 1

1

Risk factor

Treatment-specific

Myeloma-Specific

Patient or myeloma-specific

Patient-specific

- If ≥ 2 risk factors present, LMWH (equivalent to enoxaparin 40 mg/day) or full-dose warfarin (target INR: 2-3)
- VTE prophylaxis for myeloma therapy—related risk factors (eg, highdose dexamethasone, IMiDs, doxorubicin, multiagent chemotherapy)
 - LMWH (equivalent to enoxaparin 40 mg/day) or full-dose warfarin (target INR: 2-3)

Palumbo A, et al J Clin Oncol. 2014;32:587-600. Palumbo A, et al. Leukemia. 2008;22:414-423.

anemia

- Present in almost 75% of all newly dx pts and universally in relapsed disease
- Causes multifactorial both disease specific and treatment-induced
- Renal insufficiency common

Kyle et al: Mayo Clin Proc 2003. Willan et al: Clin Interv Aging 2016. Silvestris et al: Blood 2002. Birgegard et al: EurJH 2006.

Management of anemia

- PRBC transfusion for rapid restoration
- Erythropoiesis-stimulating agents (ESAs)(Erythropoietin and darbepoetin)
 - No clear guidelines for use in MM
 - Conflicting study results in patients with MM
 - Vista, subanalysis, revealed no inferior outcome with ESA use in MM but small subset
 - ASH and ASCO recommends ESA use at lowest possible dose at avoid transfusions atol 2008. Richardson et al: BJH 2011

ESA-use considerations

- · Important potential side-effects
 - Increase thromboembolic complications
 - HTN
 - Possible increased mortality
- Consider use in patients with
 - Persistent hgb <10 or who are transfusion dependent</p>
 - Ruled out other reversible causes such as iron/B12 def
 - Target hgb should be no more than 12 g/dL
- ESA use remains rooms to vers pd Baz et al: Acta Haematol 2007.

CArdiotoxicity

- Mechanism unclear
 - Direct endothelial effects, Sarcomeric protein turnover inhibition are hypotheses
- Thought to be a class effect for PIs
 - Cardiac failure Gr 3 or higher
 - Vd 3%, IxaRd 3%, KRd 7%, Kd 6%
 - HTN Gr 3 or higher
 - Vd 3%, IxaRd 3%, KRd 4%, Kd 9%
- Carfilzomib most concerning
 - All grade dyspnea 25%, HTN 15%,
 - In absence of other cardiac risk factors, severe cardiac failure ~5%
- Additional risk factors

Other cardiotoxic agents, mediastinal XRT, cardiac amyloid, high dose

Subedi et al: App payer 1/2014. Kumar et al: Blood 2016. Rosenthal et al: Blood Cancer J 2016. Hasinoff et al: Cardiovasc Toxicol 2016.

Moreau et al: Lancet Oncol 2011. Stewart et al: NEJM 2015. Dimopoulos et al: Lancet Oncol 2016. Moreau et al: NEJM 2016. Danhof et al: EJH 2016.

Carfilzomib Cardiac-directed management

- · Prior to starting therapy
 - Scree for cardiovascular risk factors
 - HTN and other cardiovascular issues should be optimally controlled prior to starting
- During treatment
 - Regular clinical surveillance
 - Good BP control
 - Serial echo, BNP are of uncertain benefit
- If cardiac failure develops
 - Discontinue use
 - Detailed cardiac evaluation and treatment
 - Once cardiac function restored
 - If individual risk-benefit favorable consider restarting carfilzomib
 - Consider dose reduction

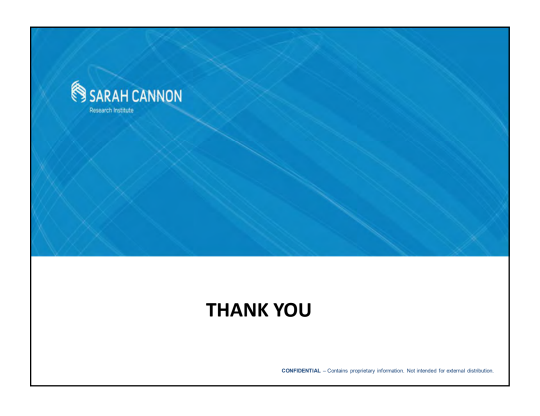
Rosenthal et al: Blood Cancer J 2016. Russell et al: Blood 2015. Mikhael: CLML 2016.

Suggested Empiric Age-Adjusted Dose Reduction in Pts With Myeloma

Agent	Younger Than 65 Yrs	65-75 Yrs	Older Than 75 Yrs
Dexamethasone	40 mg/day on Days 1-4, 15-18 q4w or Days 1, 8, 15, 22 q4w		20 mg/day on Days 1, 8, 15, 22 q4w
Melphalan	0.25 mg/kg on Days 1-4 q6w	0.25 mg/kg on Days 1-4 q6w or 0.18 mg/kg on Days 1-4 q4w	0.18 mg/kg on Days 1-4 q6w or 0.13 mg/kg on Days 1-4 q4w
Cyclophosphamide	300 mg/day on Days 1, 8, 15, 22 q4w	300 mg/day on Days 1, 8, 15 q4w or 50 mg/day on Days 1-21 q4w	50 mg/day on Days 1-21 q4w or 50 mg/day QOD on Days 1-21 q4w
Thalidomide	200 mg/day	100 mg/day or 200 mg/day	50 mg/day to 100 mg/day
Lenalidomide	25 mg/day on Days 1- 21 q4w	15-25 mg/day on Days 1- 21 q4w	10-25 mg/day on Days 1- 21 q4w
Bortezomib	1.3 mg/m² bolus on Days 1, 4, 8, 11 q3w	1.3 mg/m² bolus on Days 1, 4, 8, 11 q3w or on Days 1, 8, 15, 22 q5w	1.0- 1.3 mg/m² bolus on Days 1, 8, 15, 22 q5w

Palumbo A, et al. N Engl J Med. 2011;364:1046-1060.

emn recs)					
Complicat ion	Treatme nt	AE grade	Dosing Recommendations	Management	
Neutrope nia	Len- based	Gr 2/3 w infection or Gr 4	None if expected to improve or 25-50% dose reduction	GCSF until neutrophil recovery	
Anemia	All	Gr 2-4 (hgb <10 g/dL)	None if expected to improve or 25-50% dose reduction	Consider ESA use	
Renal	Len	CrCl 30-60 mL/min CrCl < 30 mL/min, no HD CrCl < 30 mL/min, yes HD	10 mg QD 15 mg QOD 5 mg QD	Correct other associated factors: dehydration, hypercalcemia, hyperuricemia, other nephrotoxic meds	
Neuropat hy	Thal- based	Gr 2 Gr 3-4	50% dose reduction Discontinue until Gr 1 then restart at 50% dose	Monitor carefully Consider symptomatic treatment	
	Bor- based	Gr 1 with pain or Gr 2 Gr 2 with pain or Gr 3	Switch to SQ from IV; Switch to weekly dosing Dose reduce from 1.3 to 1.0 to 0.7 mg/m2 Hold and resume when Gr 1 or less Recs as in lower grade but dose reduction rec Discontinue use	Monitor carefully Consider symptomatic treatment	
VTE	Thal/Le	All grades	Temporary discontinuation and full anticoagulation	Reevaluate for retreatment	





Thank you for attending!

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