

Sarcomas ginecológicos **leiomiosarcoma uterino**

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XV Curso Avanzado en
Sarcomas 2023



Universidad
Europea MADRID



Máster en Tumores Musculoesqueléticos

Leiomiomasarcoma <1% tumores malignos

3-10% de tumores uterinos

0,8casos/100000mujeres/a

Factores de riesgo : nuliparidad, obesidad.

Tamoxifeno y radioterapia pélvica previa.

Postmenopáusicas. Raza afroamericanas

Suelen ser esporádicos, a veces asociados a

LiFraumeni, Retinoblastoma hereditario,

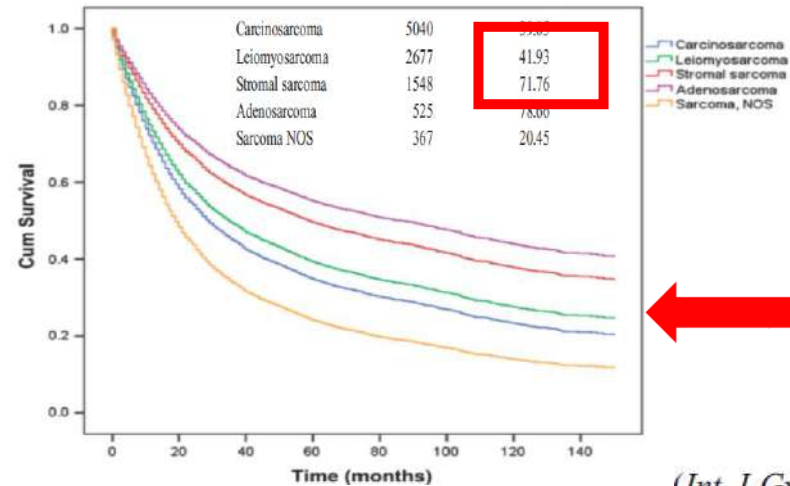
leiomiomatosis hereditaria, Mutación de la vía

germinal de la fumarato hidratasa

Clasificación WHO 2014

Estadíaaje: FIGO, AJCC

- Leiomiomasarcoma 65-70%
- Sarcoma estroma endometrial 20%
- Sarcoma del estroma endometrial de alto grado
- Sarcoma indiferenciado



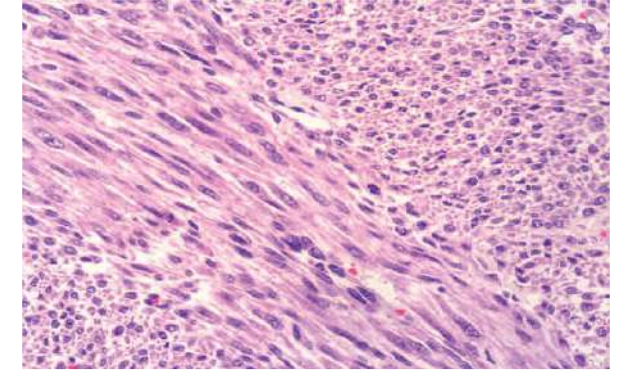
Supervivencia

(Int J Gynecol Cancer 2016;00: 00-00)

Leiomioma : clínica, dx y AP

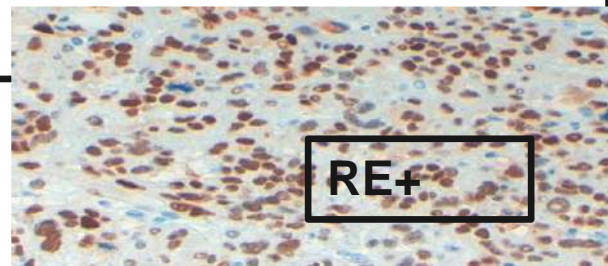
- **Clínica:** metrorragia 56%
Dolor abdominal 22%
Masa uterina 54%
- 69% localizados al diagnóstico
- 45-75% de localizados desarrollan metástasis sobre todo en los dos primeros años
- Ocasionalmente metástasis tardías

Síntomas vagos
difíciles de
diferenciar de mioma

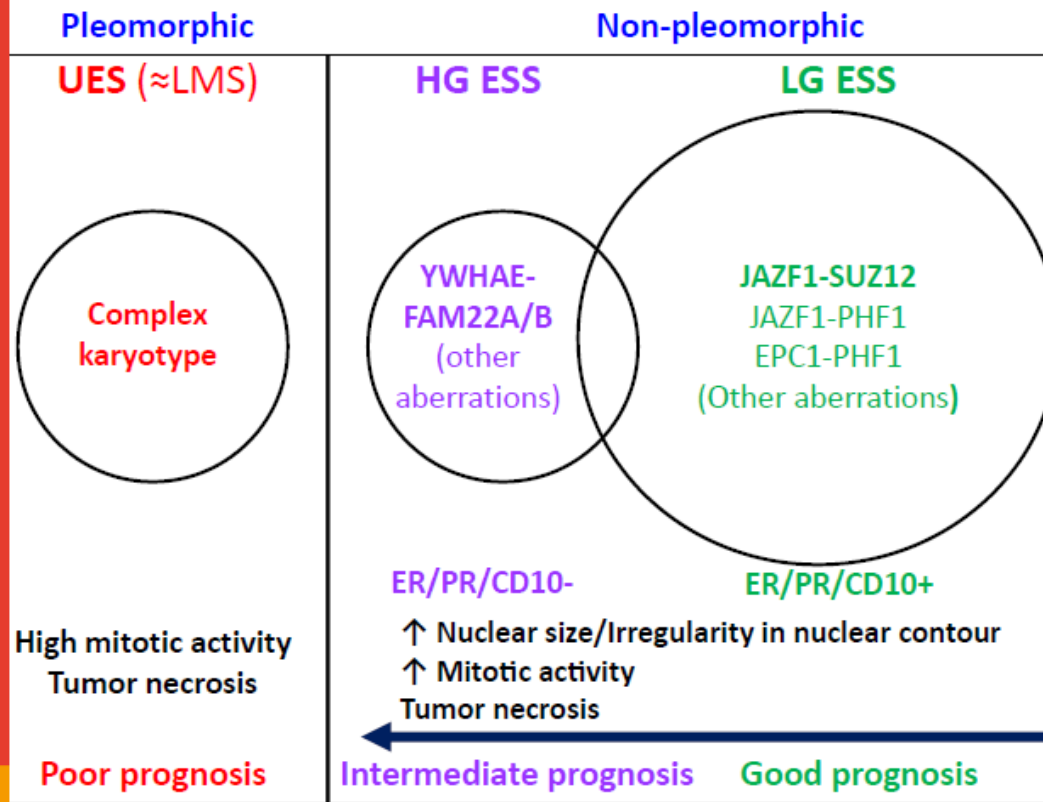


- **RX** Masa uterina con necrosis, hemorragia y puede tener calcificaciones
- A veces difícil de diferenciar de mioma. La bp preoperatoria con frecuencia negativa
- Datos de malignidad: necrosis, captación temprana de contraste, >10cm, infiltración local, crecimiento rápido, restricción a la difusión. RNM
- Metástasis: ganglios, hígado, pulmón (40%), peritoneo. Otras localizaciones más raras (cerebro, hueso...). Estudio de extensión TAC toracoabdominopelvis.

- **AP** Tumores solitarios de >5 cm con invasión a través del miometrio
- Células fusiformes con citoplasma eosinófilo
- Hiper celularidad, **atipia, necrosis, mitosis > 10M/10CGA**, hemorragia
- Desmina, caldesmon, actina y CD10 positivos
- No se gradan
- Variantes raras: leiomioma mixoide (25% fusiones PLAG1), epitelioide (algunos con reordenamiento del gen del receptor de progesterona)(puede ser queratina +)
- RE y RP positivos 30-40% casos
- Se describen casos ckit positivos (no mutaciones)



Proposed classification of uterine sarcoma



Problemas de los estudios en sarcomas uterinos

- Estudios pequeños
- Con frecuencia análisis de casos o estudios retrospectivos
- Fase II
- Mezcla de estadios
- Mezcla de localizaciones
- Mezcla de histologías
- Con frecuencia se cierran por falta de reclutamiento
- Difícil sacar conclusiones

Tratamiento. Tumor localizado leiomioma



- Cirugía: histerectomía total
- Doble anexectomía si afectación macroscópica (5%). Qué ocurre en casos con RH +?
- Afectación ganglionar rara (6-9%)
- Linfadenectomía si afectación macroscópica o afectación extrauterina
- Cirugía no planeada (histerectomía subtotal o morcellation) se aconseja nueva cirugía (upstaging 15%)
- Morcellation Aumento de riesgo de diseminación y peor pronóstico (JCO Sept 2019)
- FDA en 2014 y 2018 aconseja no realizar esta técnica
- No se aconseja cirugía para preservar fertilidad

Phase III randomised study to evaluate the role of adjuvant pelvic radiotherapy in the treatment of uterine sarcomas stages I and II An European Organisation for Research and Treatment of Cancer Gynaecological Cancer Group Study (protocol 55874)

EUROPEAN JOURNAL OF CANCER 44 (2008) 808-818

N.S. Reed^{a,c}, C. Mangioni^b, H. Malmström^c, G. Scarfone^d, A. Poveda^a, S. Pecorelli^f, S. Tateo^g, M. Franchi^h, J.J. Jobsenⁱ, C. Coens^j, I. Teodorovic^j, I. Vergote^k, J.B. Vermorken^l

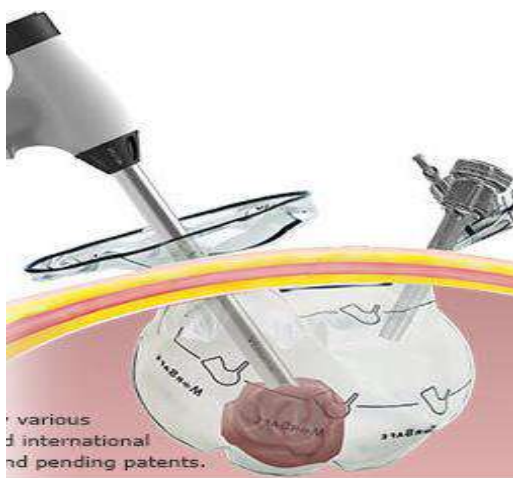
Table 6 - Sites of recurrence

	Sites of recurrence			
	CS, n = 91		LMS, n = 99	
	Radiotherapy (n = 46)	Observation (n = 45)	Radiotherapy (n = 50)	Observation (n = 49)
No local recurrence	28 (61%)	21 (47%)	22 (44%)	26 (53%)
Local recurrence only	2 (4%)	11 (24%)	1 (2%)	7 (14%)
Distant metastases	7 (15%)	3 (7%)	18 (36%)	7 (14%)
Local followed by distant	1 (2%)	3 (7%)	0 (0%)	2 (4%)
Distant followed by local	2 (4%)	0 (0%)	2 (4%)	3 (6%)
Simultaneous local and distant	6 (13%)	7 (16%)	7 (14%)	4 (8%)
Any local recurrence	11 (24%)	21 (47%)	10 (20%)	12 (24%)
Any distant metastases	16 (35%)	13 (29%)	27 (54%)	16 (33%)

50,4 Gy hasta 8 semanas tras cirugía. NO disminución de recidiva local, ni a distancia, ni aumento de supervivencia

Máster en Tumores Musculoesqueléticos

RT adyuvante Se valora en casos seleccionados: margen afecto, cirugía no reglada, afectación de parametrios, afectación cervical



histologías

	Arm A CT+RT 39	Arm B RT 42
Median follow-up	4.3 years (0.5–8.7)	4.3 years (0.3–7.9)
Relapse (patients)	15 (38.5%)	26 (62%)
3-year DFS (95% CI)	55% (40% to 70%)	41% (27% to 57%)
3-year OS (95% CI)	81% (66% to 91%)	69% (52% to 82%)
5-year OS (95% CI)	72% (53% to 85%)	55% (37% to 72%)

- Adriamicina 50mg/m², Ifos 6gr/m², Cisplatino 75mg/m² x 4-----RT Vs RT
- Se cerró por falta de reclutamiento
- 24/29 leiomiomas
- Estadios I-III

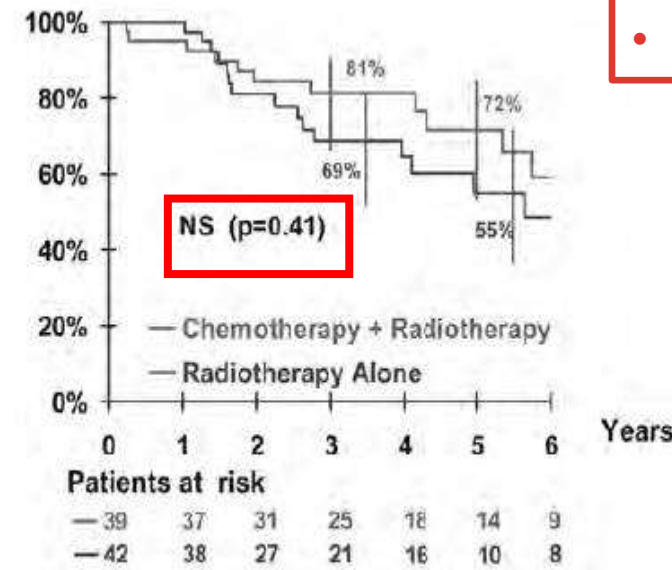
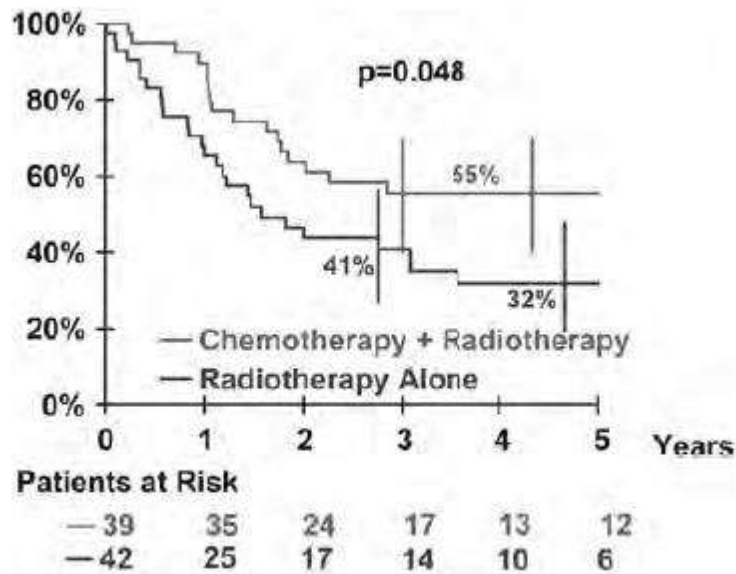


Figure 1. Disease-free survival according to adjuvant therapy group. re 2. Overall survival according to adjuvant therapy group. Vertical

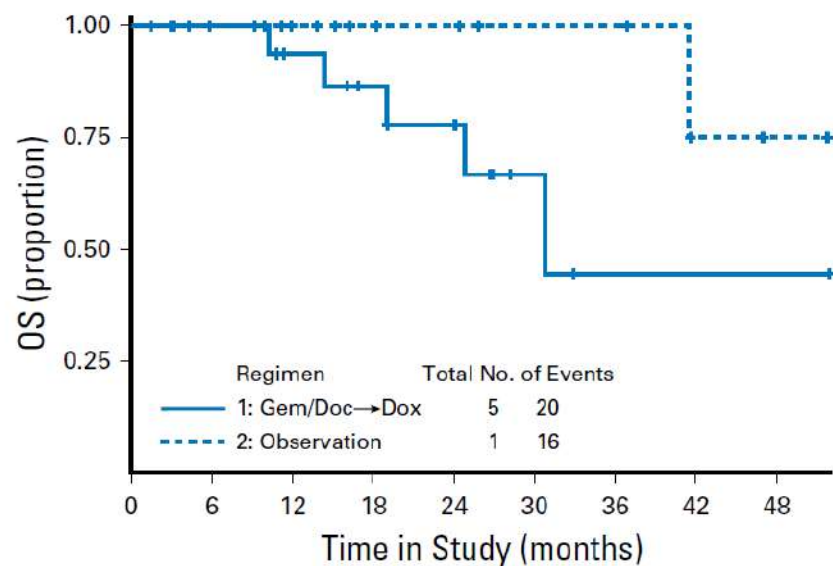
- Estudio pequeño Mezcla histologías
- No aumento de supervivencia 3% mortalidad

Adjuvant Gemcitabine Plus Docetaxel Followed by Doxorubicin Versus Observation for High-Grade Uterine Leiomyosarcoma: **A Phase III** NRG Oncology/Gynecologic Oncology Group Study

J Clin Oncol 36:3324-3330. © 2018

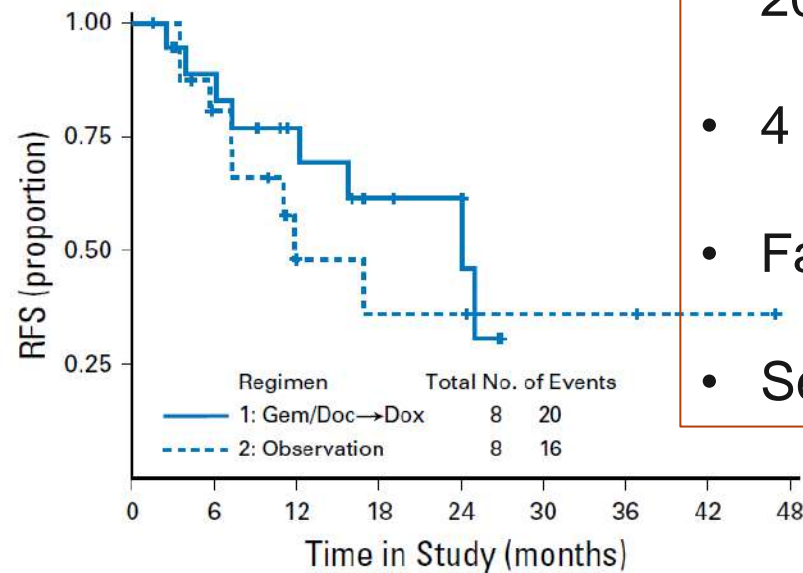
Martee L. Hensley, Danielle Enserro, Helen Hatcher, Petronella B. Ottevanger, Anders Krarup-Hansen, Jean-Yves Blay, Cyril Fisher, Katherine M. Moxley, Shashikant B. Lele, Jayanthi S. Lea, Krishmansu S. Tewari, Premal H. Thaker, Oliver Zivanovic, David M. O'Malley, Katina Robison, and David S. Miller

A



No. at risk:		0	6	12	18	24	30	36	42	48
Gem/Doc→Dox	1	20	17	13	10	8	3	1	1	1
Observation	2	16	14	11	8	7	5	5	2	1

B



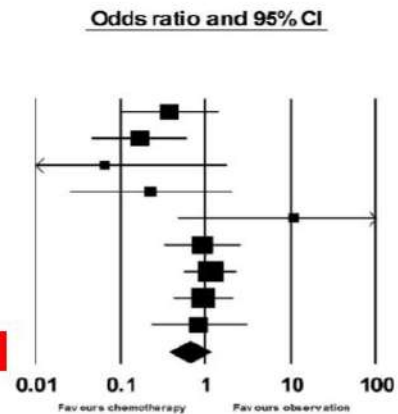
No. at risk:		0	6	12	18	24	30	36	42	48
Gem/Doc→Dox	1	20	15	10	6	5	0			
Observation	2	16	11	4	3	3	2	2	1	0

- Cerrado en Septiembre 2016
- 4 años abierto
- Falta de reclutamiento
- Se incluyeron 36 pacientes

Effect of adjuvant therapy on the risk of recurrence in early-stage leiomyosarcoma: A meta-analysis

Gynecologic Oncology 154 (2019) 638-650

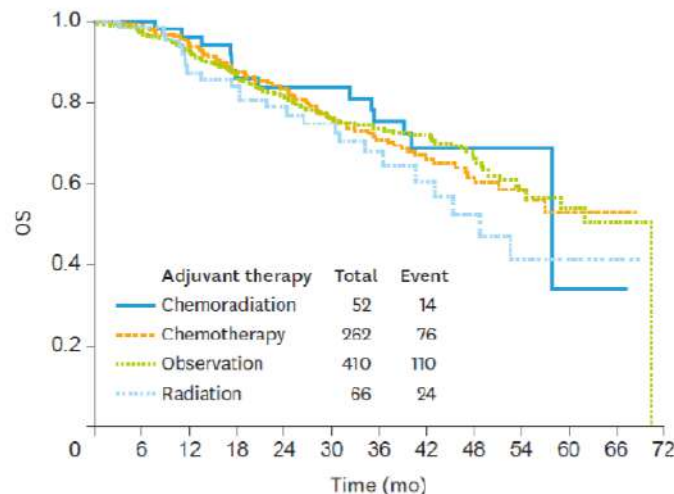
Study name	Statistics for each study				p-Value
	Odds ratio	Lower limit	Upper limit	Z-Value	
Omura, 1985	0.377	0.099	1.430	-1.435	0.151
Hornback, 1986	0.167	0.045	0.612	-2.700	0.007
Piver, 1988	0.065	0.002	1.794	-1.615	0.106
Wu, 2006	0.227	0.025	2.062	-1.317	0.188
Kim, 2009	10.818	0.463	252.789	1.481	0.139
Ricci, 2013	0.916	0.326	2.574	-0.166	0.868
Mancari, 2014	1.127	0.553	2.296	0.328	0.743
Littell, 2017	0.933	0.412	2.111	-0.167	0.868
Hensley, 2018	0.833	0.229	3.028	-0.277	0.782
	0.652	0.368	1.154	-1.470	0.142



- 747 pts/12 estudios
- Analiza QT y RT
- Cualquier recurrencia
- Se incluyen los datos de Gemcitabina TXT
- NO BENEFICIO

Characterizing the efficacy and trends of adjuvant therapy versus observation in women with early stage (uterine confined) leiomyosarcoma: a National Cancer Database study

J Gynecol Oncol. 2020 May;31(3):e21



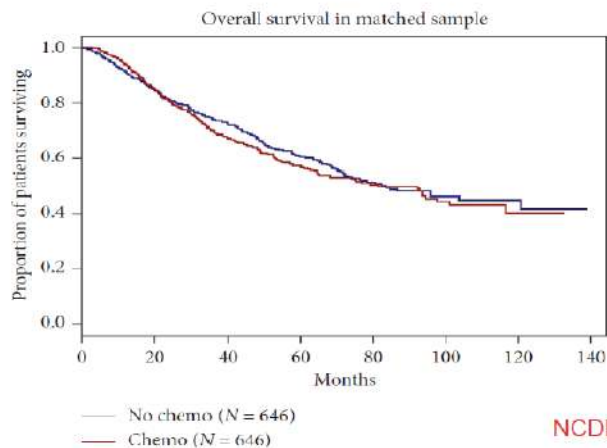
Supervivencia NO BENEFICIO

1030 mujeres
Leiomyosarcoma uterino localizado
2008-2014

Research Article

Adjuvant Chemotherapy in Uterine Leiomyosarcoma: Trends and Factors Impacting Usage

Sarcoma 2019



NO BENEFICIO

NCDB 2004-14

A randomized phase II study of letrozole vs. observation in patients with newly diagnosed uterine leiomyosarcoma (uLMS)

Brian M. Slomovitz^{a,*}, Michael C. Taub^a, Marilyn Huang^a, Charles Levenback^b,

Gynecologic Oncology Reports 27 (2019) 1-4

Enfermedad limitada al útero >10%RE
Cerrado por falta de reclutamiento

9 pacientes

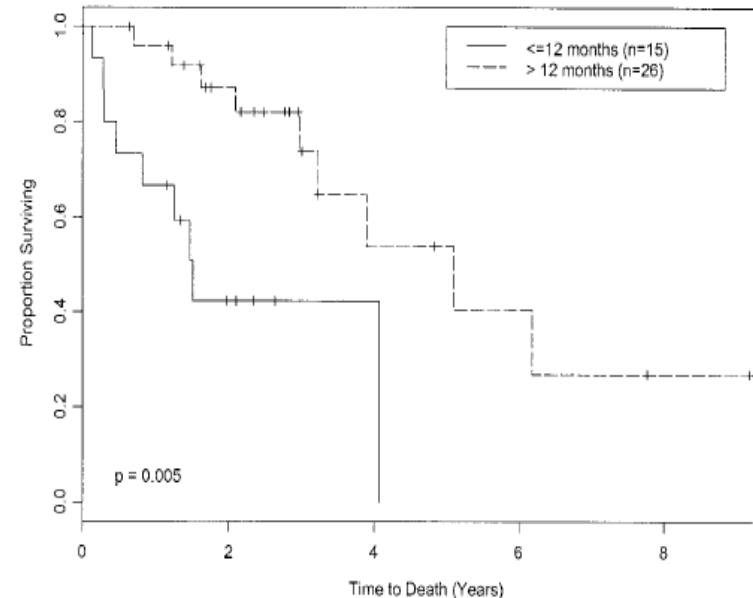
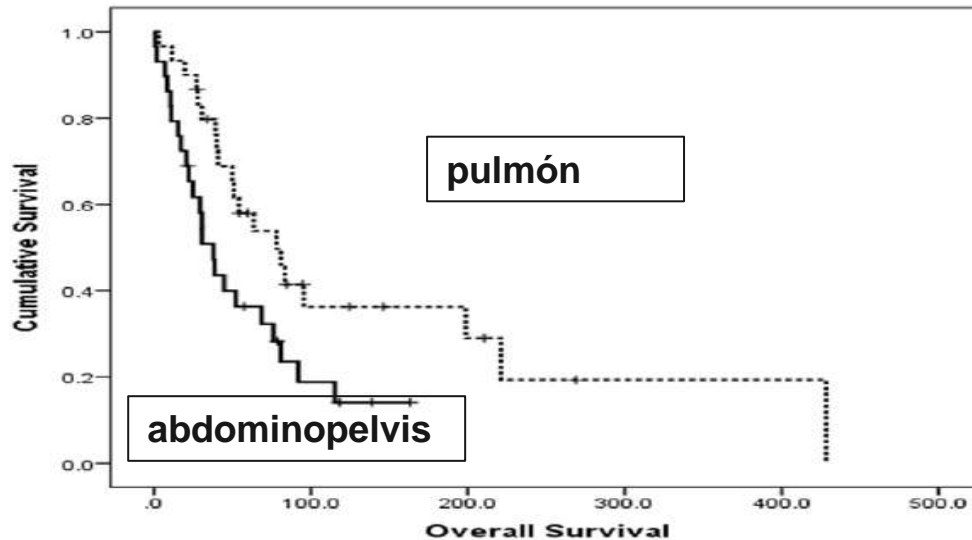
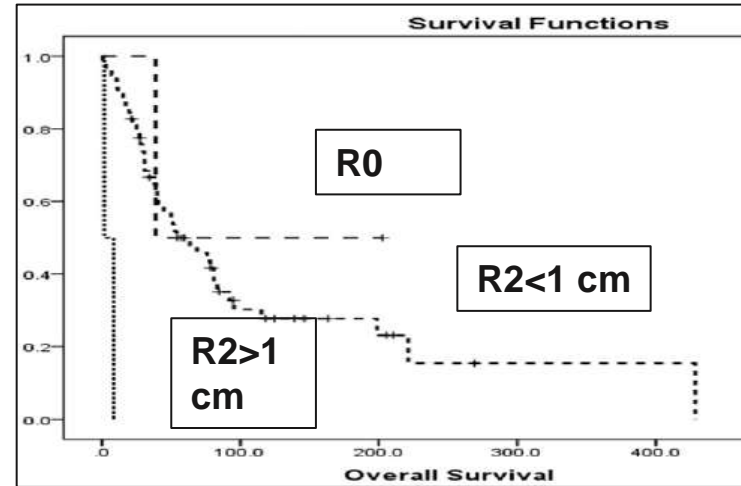
- Tratamiento estándar cirugía
- No quimioterapia ni radioterapia adyuvante salvo casos seleccionados

Secondary surgical resection for patients with recurrent uterine leiomyosarcoma

Gynecologic Oncology 154 (2019) 333–337

Mario M. Leitao Jr

- 62 pacientes
- 29 recidiva abdominopélvica
- 30 recidiva pulmonar
- 3 pulmonar y abdominopélvica
- Cirugía R0 58 p (93%)
- Supervivencia R0 54,1 m
- NO beneficio de tratamiento adyuvante



Leiomiosarcoma uterino metastásico: primera línea



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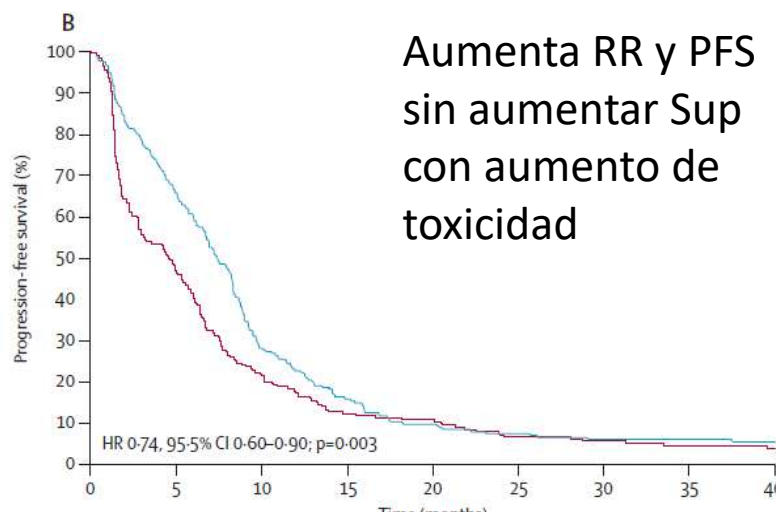
Doxorubicin alone versus intensified doxorubicin plus ifosfamide for first-line treatment of advanced or metastatic soft-tissue sarcoma: a randomised controlled phase 3 trial

Ian Judson, Jaap Verweij, Hans Gelderblom, Jörg T Hartmann, Patrick Schöffski, Jean-Yves Blay, J Martijn Kerst, Josef Sufliarsky, Jeremy Whelan,

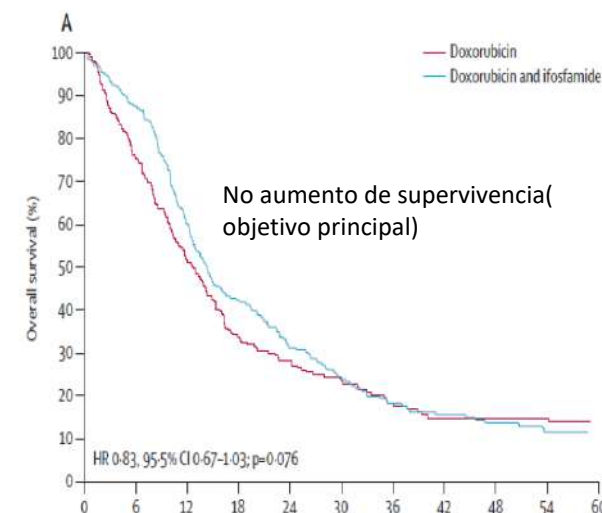
	Doxorubicin group (n=228)	Doxorubicin and ifosfamide group (n=227)
Complete response	1 (<1%)	4 (2%)
Partial response	30 (13%)	56 (25%)
Stable disease	105 (46%)	114 (50%)
Progressive disease	74 (32%)	30 (13%)
Early death (progression)	4 (2%)	5 (2%)
Early death (other cause)	3 (1%)	2 (1%)
Not evaluable	11 (5%)	16 (7%)

Data are n (%).

Table 3: Responses to treatment



Aumenta RR y PFS sin aumentar Sup con aumento de toxicidad

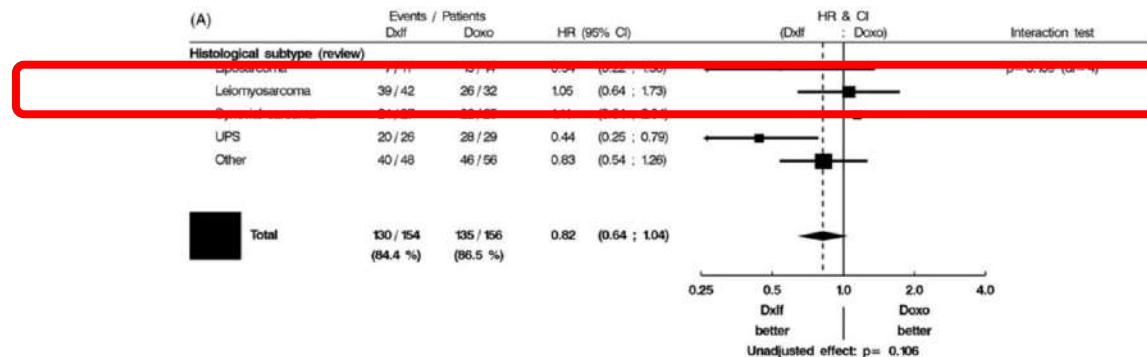


No aumento de supervivencia (objetivo principal)

	Doxorubicin group (n=228)	Doxorubicin and ifosfamide group (n=227)
Progression of disease or death caused by progressive disease	95 (42%)	47 (21%)
Toxic effect (including toxic death)	6 (3%)	40 (18%)
Toxic death	5 (2%)	2 (1%)
Patient's refusal (not related to toxic effects)	4 (2%)	10 (4%)
Intercurrent death (not related to malignant disease or toxic effects)	4 (2%)	1 (<1%)
Other	12 (5%)	11 (5%)

Data are n (%).

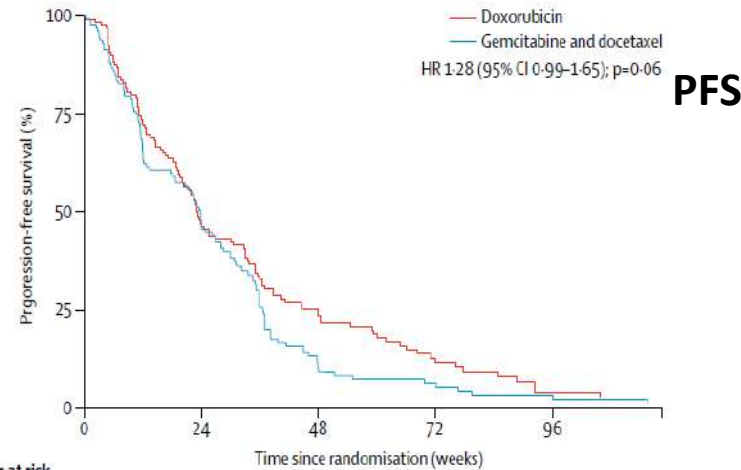
Table 4: Reasons for discontinuation of treatment



Gemcitabine and docetaxel versus doxorubicin as first-line treatment in previously untreated advanced unresectable or metastatic soft-tissue sarcomas (GeDDiS): a randomised controlled phase 3 trial

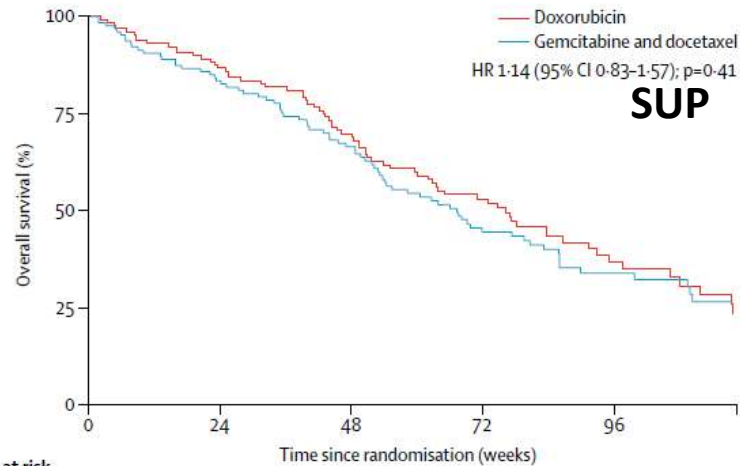
Lancet Oncol. 2017

Beatrice Seddon, Sandra J Strauss, Jeremy Whelan, Michael Leahy, Penella J Woll, Fiona Cowie, Christian Rothermundt, Zoe Wood,



Number at risk (number censored)

	0	24	48	72	96
Doxorubicin	129 (0)	60 (1)	28 (6)	11 (10)	3 (11)
Gemcitabine and docetaxel	128 (0)	60 (2)	12 (4)	5 (6)	3 (6)



Number at risk (number censored)

	0	24	48	72	96
Doxorubicin	129 (0)	108 (4)	80 (12)	47 (27)	20 (42)
Gemcitabine and docetaxel	128 (0)	104 (3)	74 (13)	44 (20)	24 (31)

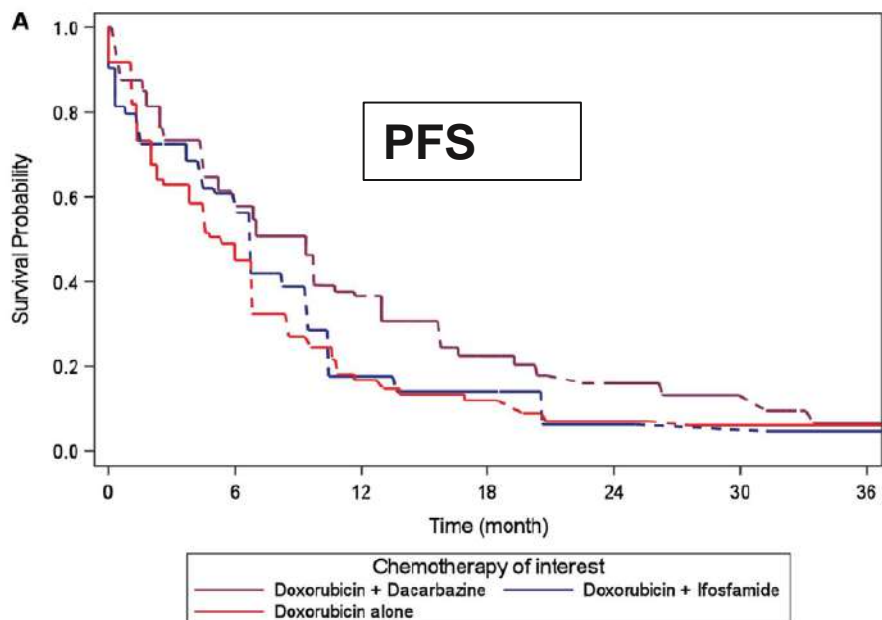
	Doxorubicin (n=129)	Gemcitabine and docetaxel (n=128)
Complete response	2 (2%)	0
Partial response	23 (18%)	25 (20%)
Stable disease	60 (47%)	50 (39%)
Progressive disease	25 (19%)	27 (21%)
Not evaluable	19 (15%)	26 (20%)

Data are n (%).

La Gemcitabina-Taxotere no aumenta la tasa de respuestas ni la supervivencia libre de progresión, ni la supervivencia con aumento de toxicidad

Doxorubicin Plus Dacarbazine, Doxorubicin Plus Ifosfamide, or Doxorubicin Alone as a First-Line Treatment for Advanced Leiomyosarcoma: A Propensity Score Matching Analysis From the European Organization for Research and Treatment of Cancer Soft Tissue and Bone Sarcoma Group

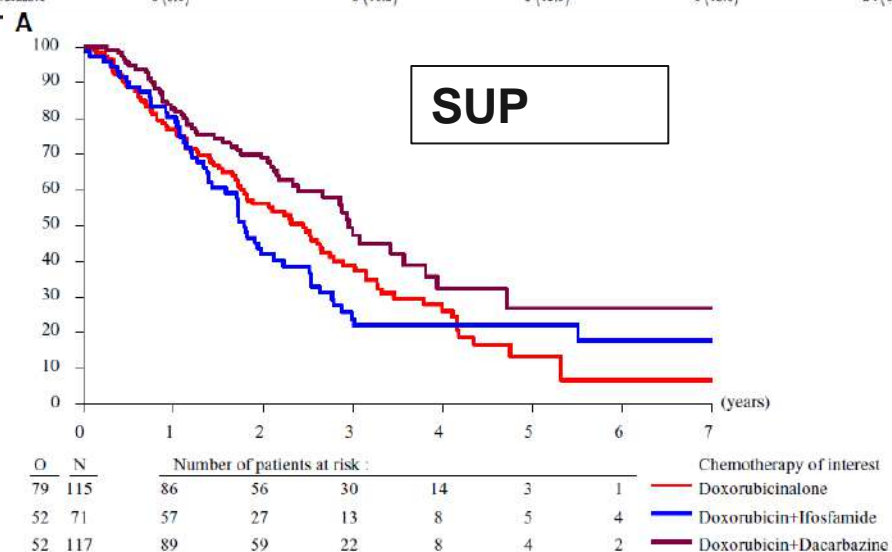
Cancer 2020;0:1-11.



RR 30,9%AD, 19,5%
AI,25,6% A
Estudio retrospectivo

Table 4
Response to chemotherapy regimen.

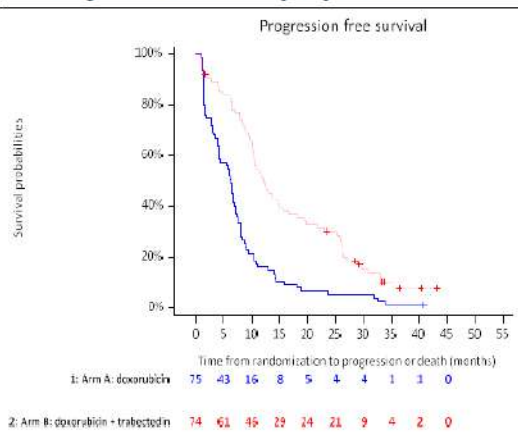
	Treatment				Total (N = 269)
	Anthracyclins (N = 119)	DOX + IFO (N = 87)	CYVADIC N = 23	IFO alone (N = 40)	
	N (%)	N (%)	N (%)	N (%)	N (%)
Best overall response					
Complete response	3 (2.5)	2 (2.3)	3 (13.0)	0 (0.0)	8 (3.0)
Partial response	26 (21.8)	19 (21.8)	5 (21.7)	2 (5.0)	52 (19.3)
No change	51 (42.9)	29 (33.3)	7 (30.4)	15 (37.5)	102 (37.9)
Progression	33 (27.7)	28 (32.2)	5 (21.7)	17 (42.5)	83 (30.9)
Non evaluable	6 (5.0)	9 (10.3)	3 (13.0)	6 (15.0)	24 (8.9)



LMS-04 study: a randomised, multicenter phase-III study comparing doxorubicin alone versus doxorubicin with trabectedin followed by trabectedin in non-progressive patients as first-line therapy in patients with metastatic or unresectable lei

LMS 04: Ph-III first-line therapy for locally advanced/metastatic LMS

PFS by BICR, ITT population



Events, n (%)

Median PFS, months

Doxo (N = 76)	Doxo + Trab (N = 74)
74 (97%)	65 (88%)
6.2	12.2
HR 0.41	
95% CI 0.29-0.58; P<0.0001	

FIRST LINE

- Ut-LMS; ST-LMS
- Locally advanced /meta
- No previous CT

Stratification

- Uterus vs soft tissue
- Locally advanced vs metastatic

N = 150

Randomization

Doxorubicin 75 mg/m² q3weeks*
Max 6 cycles
n = 76

Surgery if indicated

Doxorubicin 60 mg/m² + trabectedin 1,1 mg/m² q3 weeks*
Max 6 cycles
n = 74

Surgery if indicated

PR or SD

trabectedin 1,1 mg/m² 3h q3 weeks; until PD max 17 cycles

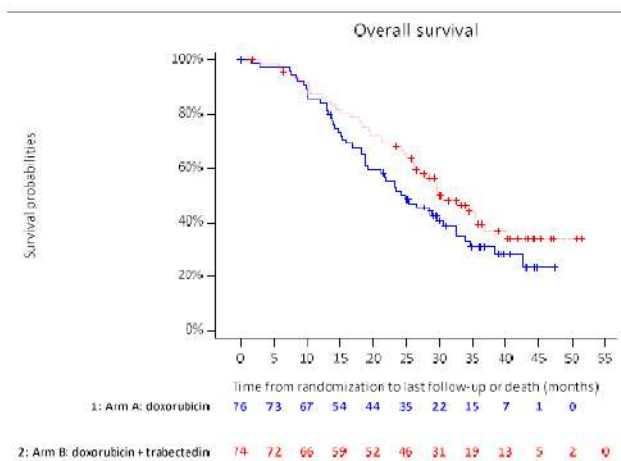
Primary endpoint
PFS (RECIST v1.1) RX review

Secondary endpoints
PFS inv
Response rate
CBR
PFS2
OS
Safety and tolerability

Efficacy

	Doxo N = 76	Doxo + Trab N = 74
Response		
> CR	0	4 (5%)
> PR	10 (13%)	24 (32%)
> SD	50 (66%)	40 (54%)
Response Rate before surgery n (%)	10 (13%)	28 (38%)
Ut-LMS (n = 67)	5 (15%)	12 (36%)
ST-LMS (n = 83)	5 (12%)	16 (39%)
CBR (CR + PR + SD)	60 (79%)	68 (92%)
Duration of response (months) Median [IQR]	5.6 [4.1-6.9]	12.5 [7.8-20.3]

Overall survival



Deaths, n (%)

Median OS, months

Doxo (N = 76)	Doxo + Trab (N = 74)
50 (66%)	42 (57%)
24.1	30.5
HR 0.73	
95% CI: 0.49-1.12	

Median follow-up : 37 months

CI, confidence interval; HR, hazard ratio; PFS, progression-free survival; Doxo: Doxorubicin; Trab: Trabectedin.

Most common Gr 3-4 AEs

Gr 3-4 AEs n (%)	Doxo N = 76	Doxo + Trab N = 74
Number pts with at least 1 Gr3-4 AEs	20 (26%)	35 (47%)
Fatigue	7 (9%)	8 (11%)
Anemia	1 (1%)	10 (14%)
Neutropenia	5 (7%)	32 (43%)
Febrile neutropenia	8 (11%)	18 (24%)
Thrombocytopenia	0	15 (20%)
Transaminase elevation	0	13 (18%)
Nausea/Vomiting	1 (1%)	9 (12%)*
Cardiac failure	2 (3%)	1 (1%)
Toxic death	1 (1%)	0



Leiomiomasarcoma uterino metastásico pretratado



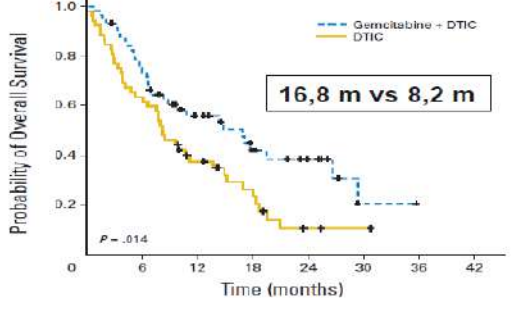
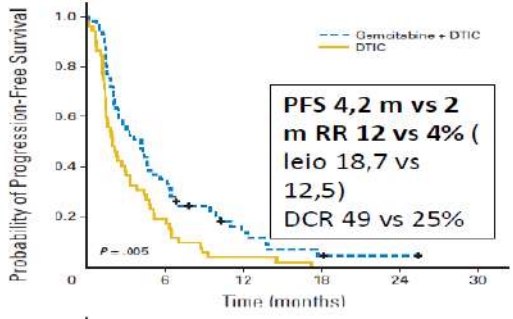
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Randomized Phase II Study Comparing Gemcitabine Plus Dacarbazine Versus Dacarbazine Alone in Patients With Previously Treated Soft Tissue Sarcoma: A Spanish Group for Research on Sarcomas Study

Xavier Garcia-del-Muro, Antonio Lopez-Pousa, Juan Mascaró, Javier Martínez, Javier Martínez-Trufero, Antonio Casado, Anacleto Gómez, Espasa, Joaquín Eiro, José María Pérez, Andrés Muñoz, Carlos Bény, Ricardo Calvo, José Andrés, Ana de Juan, María Latorre, Juan Antonio Carrasco, Raquel de Andrés, and José M. Lacort

Characteristic	DTIC (n = 52)		Gemcitabine + DTIC (n = 57)	
	No.	%	No.	%
Histologic diagnosis				
Leiomyosarcoma	16	31	16	28
Liposarcoma	9	17	10	18
Undifferentiated pleomorphic	8	15	11	19
Synovial sarcoma	5	10	6	11
Miscellaneous sarcoma	14	27	14	25
Histologic grade				
Low	5	10	4	7
High	41	79	48	84
Unknown	6	12	5	9
Site of primary				
Extremity and trunk wall	27	52	23	40
Retroperitoneum	6	12	11	19
Gynecologic	10	19	5	9
Other	9	17	18	32

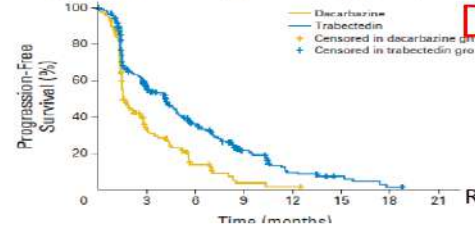


Efficacy and Safety of Trabectedin or Dacarbazine for Metastatic Liposarcoma or Leiomyosarcoma After Failure of Conventional Chemotherapy: Results of a Phase III Randomized Multicenter Clinical Trial JCO 2015

George D. Demetri, Margaret von Mehren, Robin L. Jones, Martee L. Hensley, Scott M. Schuetz

Table 1. Baseline Demographic and Disease Characteristics

Variable	Trabectedin (n = 245)	Dacarbazine (n = 178)
Age, years		
Median (range)	57 (18-81.0)	56 (17.0-79.6)
Sex		
Male	107 (21)	47 (27)
Female	238 (99)	126 (73)
Baseline BMI, kg/m ²		
Median (range)	28.21 (14.5-78.1)	27.05 (13.2-66.7)
Histology		
Leiomyosarcoma	257 (72)	176 (72)
Uterine	134 (39)	78 (45)
Nonuterine	118 (34)	48 (28)
Liposarcoma	95 (27)	47 (27)
Myxoid ± round cell	36 (11)	19 (11)
Pleomorphic	10 (3)	8 (5)
Dedifferentiated	45 (13)	25 (15)



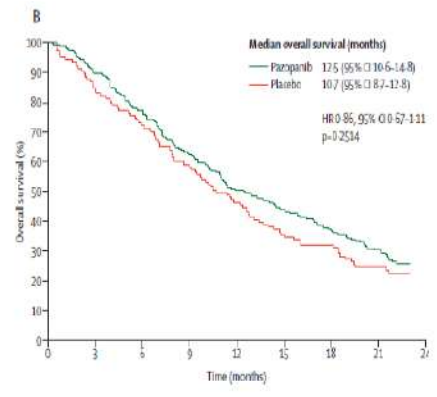
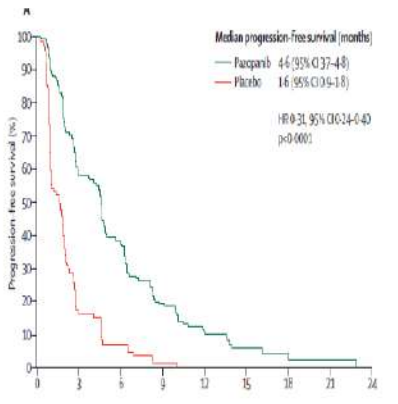
Subgroup	Median (months)		HR
	Dacarbazine	Trabectedin	
All	1.5	4.2	0.55
1	2.7	4.9	0.49
≥ 2	1.5	4.2	0.56
0	1.5	4.7	0.51
1	1.5	2.9	0.60
Leiomyosarcoma	1.6	4.3	0.55
Nonuterine	1.6	4.9	0.58
Uterine	1.5	4.0	0.58
Liposarcoma	1.5	3.0	0.55
Dedifferentiated	1.9	2.2	0.68
Myxoid ± round cell	1.5	5.6	0.41
Pleomorphic	1.4	1.5	0.33

Aumento de PFS 1,5 m vs 4,2 m sin aumento en sup

RR 9,9% vs 6,9% (beneficio clínico 34 vs 18%)
Máster en Tumores Musculoqueléticos

Pazopanib for metastatic soft-tissue sarcoma (PALETTE): a randomised, double-blind, placebo-controlled phase 3 trial

Winette TA van der Graaf, Jean-Yves Blay, Sant P Chowik, Dong-Wan Kim, Binh-Bui-Nguyen, Paolo G Cossoli, Patrick Schöffski, Massimo Aglietta, Lancet 2012; 379: 1879-86



123pts placebo/246 pts pazopanib
Objetivo principal

Randomized Multicenter and Stratified Phase II Study of Gemcitabine Alone Versus Gemcitabine and Docetaxel in Patients with Metastatic or Relapsed Leiomyosarcomas: A Fédération Nationale des Centres de Lutte Contre le Cancer (FNCLCC) French Sarcoma Group Study (TAXOGEM study)

PATRICIA PAUTIER,^a ANNE FLOQUET,^c NICOLAS PENEL,^d SOPHIE PIPERNO-NEUMANN,^e

The Oncologist 2012;17:1213-1220

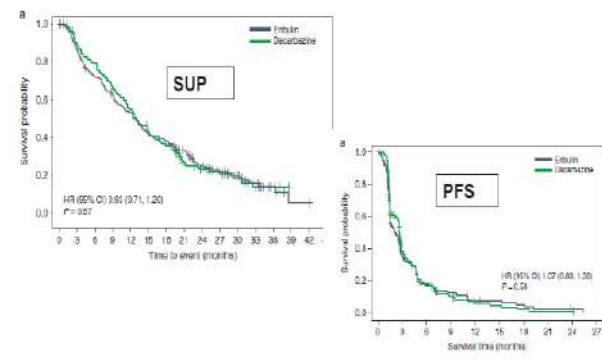
2ª línea tras antraciclina

Table 4. Responses to treatment

	Uterine group (%)	
	Gemcitabine	Gemcitabine + docetaxel
n	21	21
Assessable patients	21	21
Complete response, n (%)	1 (5)	0
Partial response, n (%)	3 (14)	5 (24)
Stable disease	9 (43)	10 (48)
Progression	8 (38)	6 (28)
Objective response, % (95% CI)	19 (5-42)	24 (8-47)
Nonprogression rate, % (95% CI)	62 (38-82)	71 (48-89)
Progression-free survival, % (95% CI)		
3 mos	57 (37-76)	71 (50-86)
6 mos	48 (28-68)	48 (28-68)
Median progression-free survival (mos)	5.5	4.7
Median overall survival (mos)	20	23

Eribulin versus dacarbazine in patients with leiomyosarcoma: subgroup analysis from a phase 3, open-label, randomised study

British Journal of Cancer (2019) 120:1026-1032; f



usculoqueléticos

A Phase II Trial of Temozolomide as a 6-Week, Continuous, Oral Schedule in Patients with Advanced Soft Tissue Sarcoma

Cancer 2005;104:1706-12.

A Study by the Spanish Group for Research on Sarcomas

- Temodal 75-100 mg/m²/d x 6 semanas/9 semanas
- 45 pts
- 7RP (15,5%)
- PFS 2,2 m
- Sup 8,1 m
- 5/11 leiomiomas ginecológicos responden
- 2 pts SLP > 3 años

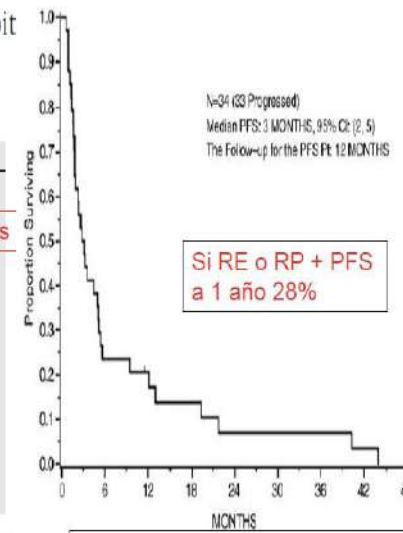
Treatment of advanced uterine leiomyosarcoma with aromatase inhibitor

Roisin O'Ceirbhail^a, Qin Zhou^b, Alexia Iasonos^b, Robert A. Soslow^c, Mario. M. Leitao^d, Carol Aghajanian^a, Martee L. Hensley^{a,*}

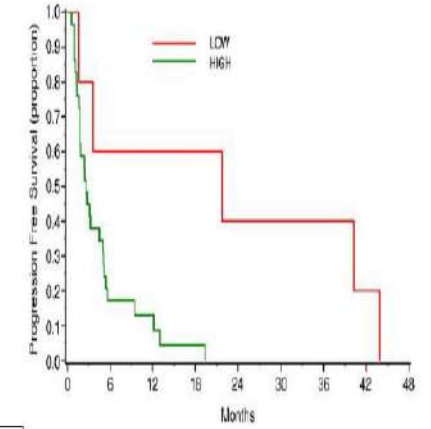
Gynecologic Oncology 116 (2010) 424-429

Variable	n (%), median (range)
Initial management at diagnosis of uLMS	
Surgical resection alone	22 (65%)
Surgical resection and chemotherapy	11 (33%)
Chemotherapy alone	1 (3%)
Number of prior chemotherapy regimens	
0	11 (32%)
1	10 (29%)
2-3	11 (32%)
≥4	2 (6%)
Prior hormonal treatment (medroxyprogesterone, tamoxifen)	7 (21%)
Prior pelvic radiotherapy	12 (35%)
Median interval between diagnosis and AI initiation	1.2 years (0.02-22)
AI used	
Letrozole [with leuprolide]	25 (74%) [3 (9%)]
Anastrozole	7 (21%)
Exemestane	2 (6%)

Response	n (%)
Complete response	0 (0%)
Partial response	3 (9%)
Stable disease	11 (32%)
Progressive disease	20 (59%)



Valorar en escaso volumen de enfermedad, lento crecimiento, R+



Máster en Tumores Musculoesqueléticos

Thanopoulos et al. Clinical Sarcoma Research 2014, 4:5
http://www.clinicalsarcomaresearch.com/content/4/1/5



RESEARCH

Open

Treatment of hormone positive uterine leiomyosarcoma with aromatase inhibitors

Eirini Thanopoulos^a, Khin Thway, Komel Khabra and Ian Judson

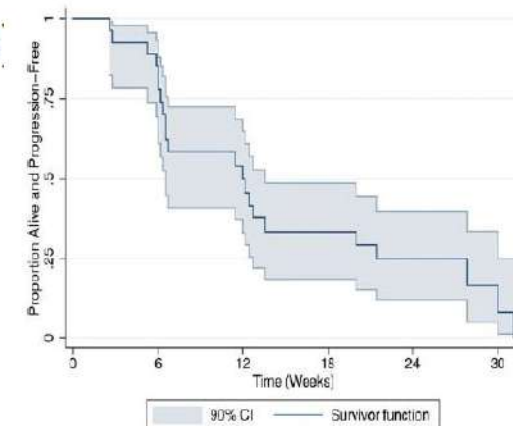
16 pacientes RE/RP +
Letrozol
PFS14 m Bajo grado 20m vs 11m.RE
+++ 20m vs 12m
RR 12,5% CBR 62%

volumen de enfermedad y lento crecimiento

Phase 2 Trial of Aromatase Inhibition With Letrozole in Patients With Uterine Leiomyosarcomas Expressing Estrogen and/or Progesterone Receptors

Cancer 2014;120:738-43

- 27 pacientes
- Mediana 2 líneas previas (0-9)
- EE 54%
- PFS 12 s
- 3 pts >24 s



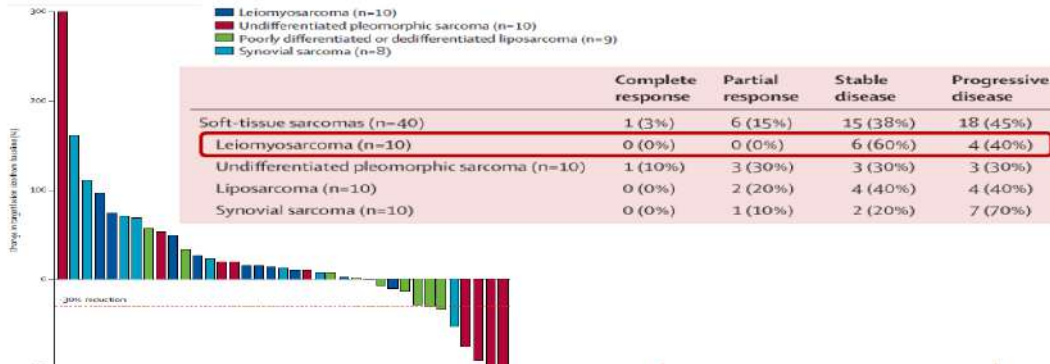
Máster en Tumores Musculoesqueléticos

population	n (%u)LMS	RR (%)	3mo PFR (%)	6mo PFR (%)	PFS (months)	OS (months)	
nivolumab (Ben-Ami et al., 2017)	uLMS	12 PDL1 + 20%	0	-	-	1.8	-
pazopanib (Benson et al., 2016)	US	44	11	-	-	3	17.5
regorafenib vs. placebo (Mir et al., 2016)	STS, 1 cohort LMS	56 LMS of who 22 uLMS	0	57 vs. 25	21 vs. 7	3.7 vs. 1.8	21 vs. 9.1
sunitinib (Hensley et al., 2009b)	uLMS	23	8.7	-	17.4	1.5	15.1
thalidomide (McMeekin et al., 2007)	uLMS	30	0	-	-	1.9	8.3
afibercept (Mackay et al., 2012)	uLMS	41	0	-	17	1.8	18.1
alisertib (Hyman et al., 2017)	uLMS	21	0	-	0	1.7	14.5

Pembrolizumab in advanced soft-tissue sarcoma and bone sarcoma (SARC028): a multicentre, two-cohort, single-arm, open-label, phase 2 trial

Hussein A Tavbi, Melissa Burgess, Vanessa Bolejack, Brian A Van Tine, Scott M Schuetz, James Hu, Sandra D'Angelo, Steven Arzoo, Richard F Riedel, Dennis A Pribaz, Sujana Mavva, Lani F Davis, Scott H Okuno, Damien R Reed, John Crowley, Lisa H Butterfield, Ruth Salazar, Jaime Rodriguez-Corales, Alexander J Lazar, Ignacio F M Stuba, Laurence H Baker, Robert G Maki, Denise Reinke, Shreyas Kumar Patel

Lancet Oncol 2017; 18: 1493-150.



- Fase III de Adriamicina- trabectedina vs adriamicina seguida de trabectedina NCT02997358
- Fase II de nivolumab en leiomiosarcoma uterino, SEE y sarcoma indiferenciado NCT03241745
- Fase II Pembrolizumab en sarcomas de partes blandas y hueso NCT02301039
- Fase I-II pembrolizumab+ Adriamicina en sarcomas de partes blandas NCT02888665
- Fase II pembrolizumab- TVEC NCT 03069378
- Neoadyuvancia con Durvalumab+ Tremelimumab + RT en sarcomas de alto riesgo NCT03116529
- Pembrolizumab+ axitinib en leiomiosarcoma NCT02636725
- Fase I-II para sarcomas avanzados primera línea Trabectedina+ Nivolumab+ Ipilimumab NCT03138161
- Fase II randomizado Pazopanib+ Gemcitabina vs Gemcitabina taxotere en sarcomas de partes blandas NCT01593748
- Gemcitabina Taxotere+ Olaratumab Fase Ib-II en sarcomas de partes blandas NCT02659020
- Inhibidores PARP en BRCA2

Assessment of Safety and Efficacy of Combined Trabectedin and Low-Dose Radiotherapy for Patients With Metastatic Soft-Tissue Sarcomas: A Nonrandomized Phase 1/2 Clinical Trial

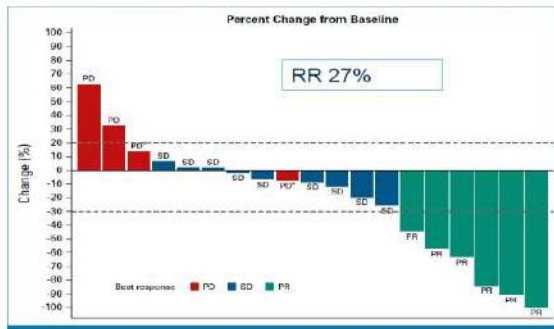
JAMA Oncol. 2020;16(2):e190004. doi:10.1001/jamaoncol.2019.6584. Published online February 20, 2020.

Javier Martin-Broto, MD, PhD; Nadia Hindi, MD; Antonio Lopez-Pousa, MD; Javier Peinado-Serrano, MD; Rosa Alvarez, MD;

Phase II Study of Olaparib and Temozolomide for Advanced Uterine Leiomyosarcoma (NCI Protocol 10250)

Matthew Ingber, MD¹, Jacob B. Alire, MS², Li Chen, PhD³, Kiewonji Dan, PhD⁴, Rose Koelgeskorok, PhD⁵, Katherine Gane, BS⁶

JCO 2023

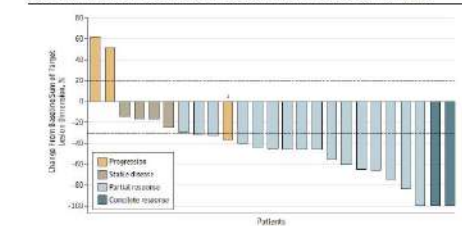


Prior lines of treatment	n (%)
1-2	9 (41)
3 or greater	13 (59)
Prior treatment received	n (%)
Gemcitabine/docetaxel	19 (86)
Doxorubicin (monotherapy)	8 (36)
Doxorubicin/olaparib	5 (23)
Doxorubicin/ifosfamide	4 (18)
Hormone therapy	6 (27)
Trabectedin	5 (23)
Investigational agent	4 (18)
Pazopanib	3 (14)

Findings In this nonrandomized phase 1/2 clinical trial of 45 patients, the maximum tolerated dose of trabectedin when combined with the specified radiotherapy regimen in phase 1 was 1.5 mg/m², which was the recommended dose used in phase 2. The overall response rate in the 25 patients with evaluable data was 72% and 60% in local and central assessments, respectively.

- Seguimiento mediana 14 meses
- PFS 9,9 m

Figure 2. Waterfall Plot of Phase 2 Trabectedin and Radiotherapy in Soft-Tissue Sarcoma Trial Results



LOCALIZADO:

- Histerectomía, no tratamiento adyuvante

PRIMERA LINEA:

- MTS La sustitución de adriamicina por otros fármacos no mejora los resultados
- La combinación de adriamicina con trabectedina aumenta la tasa de respuestas y la PFS respecto a adriamicina sola en un estudio fase III
- La combinación de adriamicina con DTIC mejora los resultados de adriamicina sola en un estudio retrospectivo
- 2as líneas Gem, Gem TXT, Gem DTIC, Trabectedina, Pazopanib.
- Mejor secuencia desconocida
- Ocasionalmente hormonoterapia
- Inclusión en ensayos clínicos



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