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8-10 września 2022 r., Bydgoszcz



Primary Results From the Double-Blind, Placebo-Controlled, Phase III SHINE Study of Ibrutinib in Combination With Bendamustine-Rituximab and Rituximab Maintenance as a First-Line Treatment for Older Patients With Mantle Cell Lymphoma

Wojciech Jurczak



BR as First-line MCL Treatment in Older Patients

- Older patients with newly diagnosed mantle cell lymphoma (MCL) are usually treated with chemo-immunotherapy regimens such as bendamustine-rituximab (BR), R-CHOP, or VR-CAP¹⁻⁴
 - BR has become the most commonly used first-line regimen⁵
- BR alone:
 - Improved progression-free survival (PFS) compared with R-CHOP (35 vs 22 months)⁶ and has a better safety profile^{6,7}
- BR with rituximab maintenance:
 - Significantly improved PFS compared with BR alone in 2 independent real world studies^{5,8}

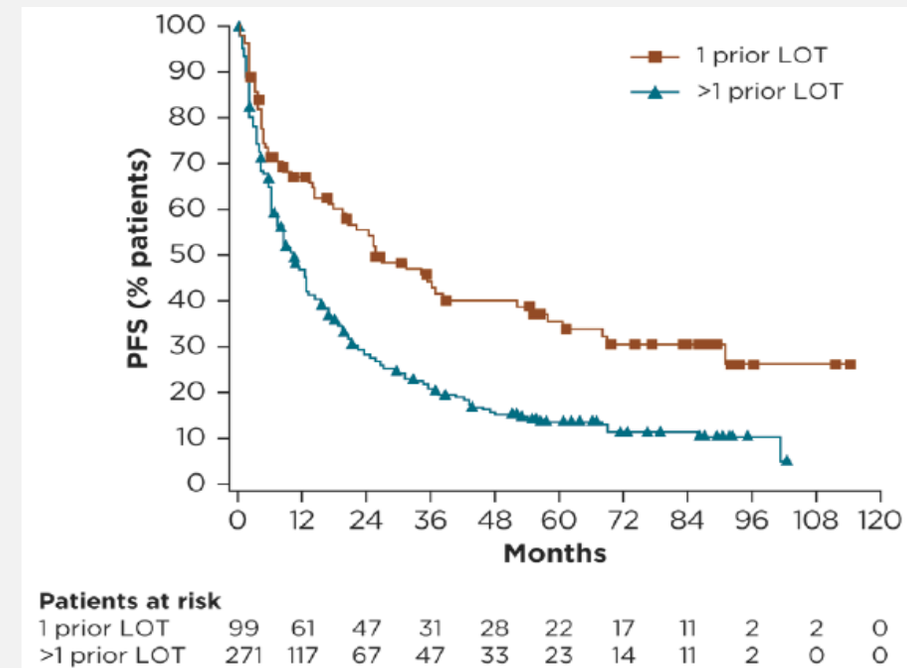
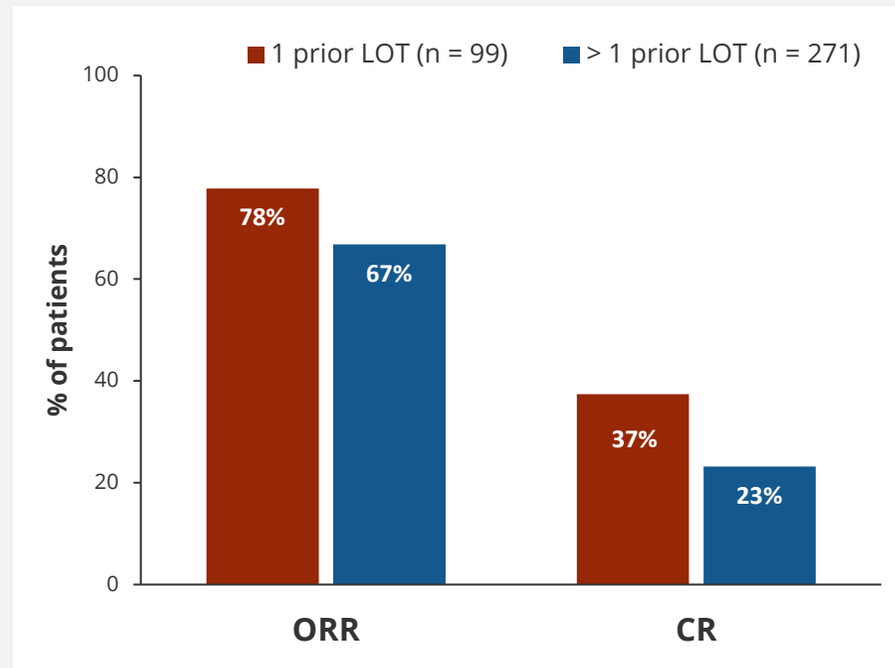
R-CHOP, rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone; VR-CAP, bortezomib, rituximab, cyclophosphamide, doxorubicin, and prednisone.

1. Hermine O, et al. *Lancet*. 2016;388:565-575. 2. Le Gouill S, et al. *N Engl J Med*. 2017;377:1250-1260. 3. Robak T, et al. *Leuk Lymphoma*. 2019;60:2622-2634. 4. Monga N, et al. *Crit Rev Oncol Hematol*. 2021;158:103212. 5. Martin P, et al. *J Clin Oncol*. 2021;39(suppl 15):7504. 6. Rummel MJ, et al. *Lancet*. 2013;381:1203-1210. 7. Flinn IW, et al. *J Clin Oncol*. 2019;37:984-991. 8. Hill BT, et al. *Hematol Oncol*. 2019;37:405-407.



Ibrutinib Is a First-in-Class Once-Daily BTK Inhibitor

- Ibrutinib has transformed the care of patients with relapsed/refractory MCL; it is particularly effective and durable at first relapse¹⁻⁵



- Ibrutinib + BR has demonstrated activity in first-line MCL in a phase 1b study⁶

BTK, Bruton's tyrosine kinase; LOT, line of therapy.

1. Wang ML, et al. *N Engl J Med*. 2013;369:507-516. 2. Rule S, et al. *Leukemia*. 2018;32:1799-1803. 3. Rule S, et al. *Blood*. 2019;134(suppl 1):1538. 4. Rule S, et al. *Haematologica*. 2019;104:e214. 5. Dreyling M, et al. *HemaSphere*. 2022;6:e712. 6. Maddocks K, et al. *Blood*. 2015;125:242-248.



SHINE: A Randomized, Double-Blind, Phase III Study

Patients

- Previously untreated MCL
- ≥ 65 years of age
- Stage II-IV disease
- No planned stem cell transplant

Stratification factor

- Simplified MIPI score (low vs intermediate vs high)

Enrolled between May 2013 and November 2014 at 183 sites

N = 523

R
1:1

BR induction for 6 cycles

if CR or PR

Rituximab maintenance every 8 weeks for 12 cycles

Ibrutinib 560 mg (4 capsules daily) until PD or unacceptable toxicity

BR induction for 6 cycles

if CR or PR

Rituximab maintenance every 8 weeks for 12 cycles

Placebo (4 capsules daily) until PD or unacceptable toxicity

Primary end point: PFS (investigator-assessed) in the ITT population

Key secondary end points: response rate, time to next treatment, overall survival, safety

Induction: Bendamustine 90 mg/m² Days 1 and 2, Rituximab 375 mg/m² Day 1, Q4W. A cycle is defined as 28 days.

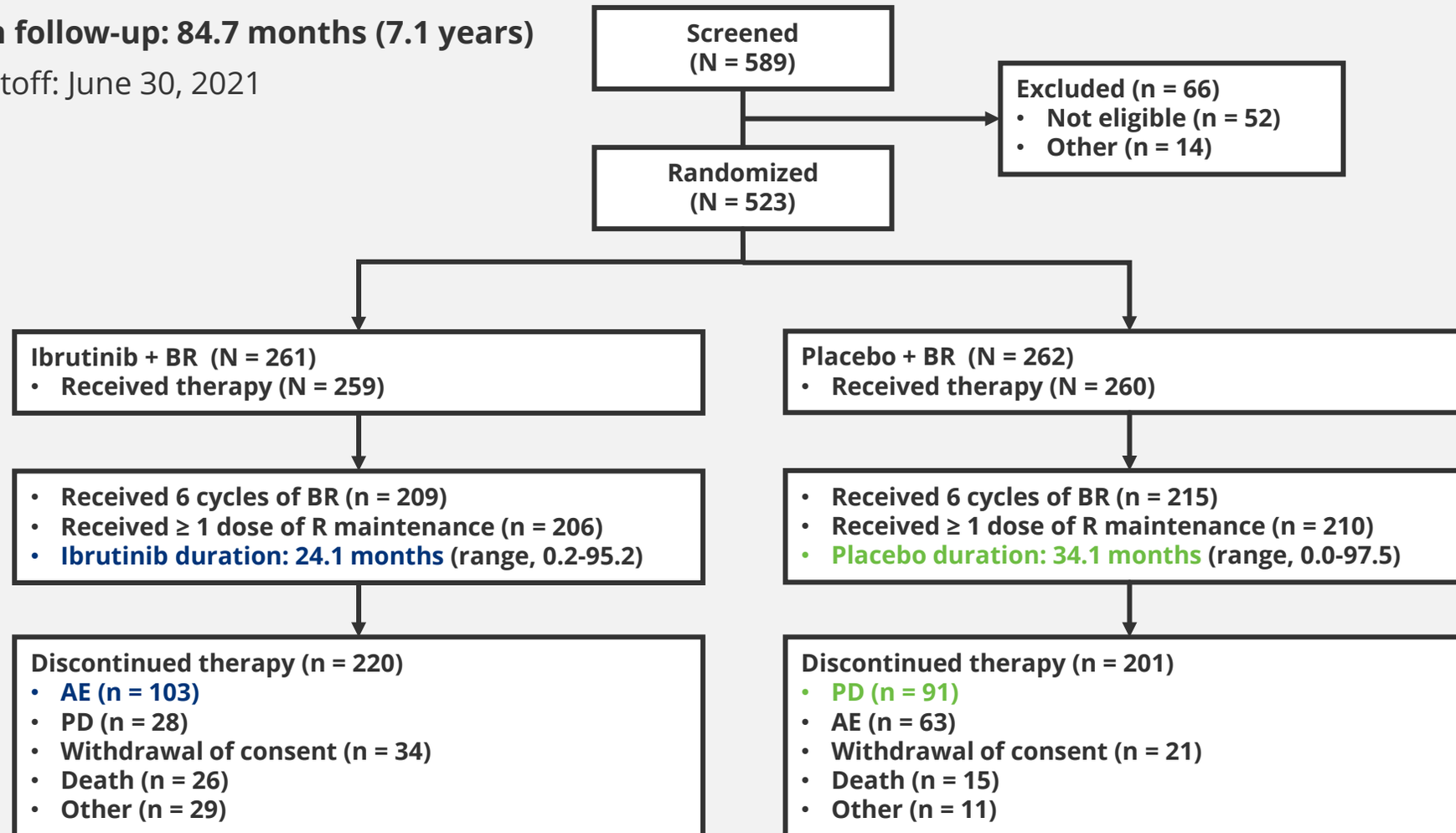
CR, complete response; ITT, intent-to-treat; MIPI, Mantle Cell Lymphoma International Prognostic Index; PD, progressive disease; PFS, progression-free survival; PR, partial response.



Patient Disposition and Treatment Exposure

Median follow-up: 84.7 months (7.1 years)

Data cutoff: June 30, 2021

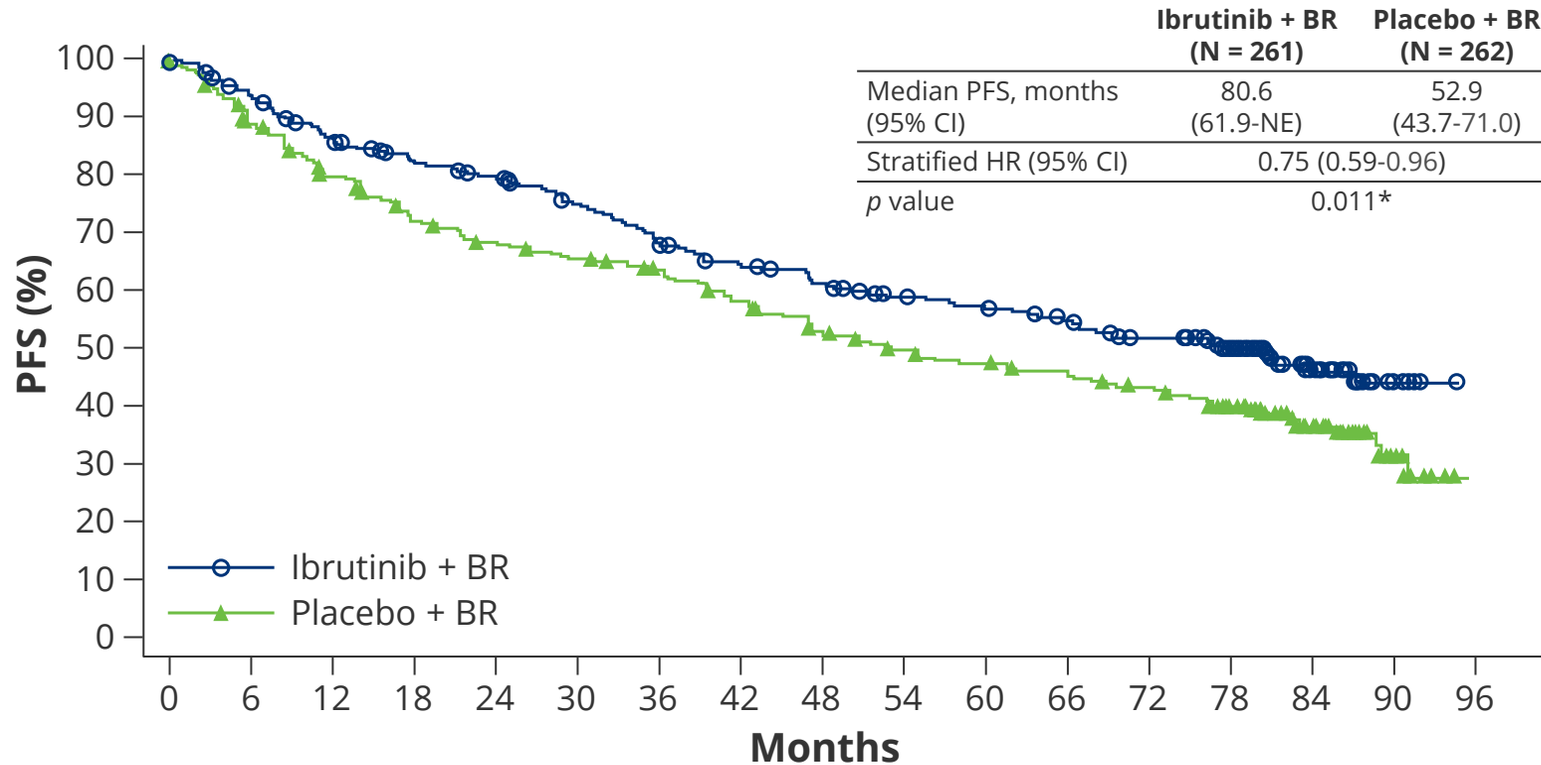


Baseline Characteristics

		Ibrutinib + BR (N = 261)	Placebo + BR (N = 262)
Median age (range), years		71 (65-86)	71 (65-87)
≥ 75 years, n (%)		74 (28.4)	82 (31.3)
Male, n (%)		178 (68.2)	186 (71.0)
ECOG PS 1, n (%)		127 (48.7)	118 (45.0)
Simplified MIPI, n (%)	Low risk	44 (16.9)	46 (17.6)
	Intermediate risk	124 (47.5)	129 (49.2)
	High risk	93 (35.6)	87 (33.2)
Bone marrow involvement, n (%)		198 (75.9)	200 (76.3)
Blastoid/pleomorphic histology, n (%)		19 (7.3)	26 (9.9)
Extranodal, n (%)		234 (89.7)	226 (86.3)
Bulky (≥ 5 cm), n (%)		95 (36.4)	98 (37.4)
TP53 mutated, n (%)		26 (10.0)	24 (9.2)
TP53 mutation status unknown, n (%)		121 (46.4)	133 (50.8)



Primary End Point of Improved PFS Was Met



Ibrutinib + BR and R maintenance achieved:

- **Significant improvement in median PFS by 2.3 years (6.7 vs 4.4 years)**
- **25% reduction** in risk of PD or death

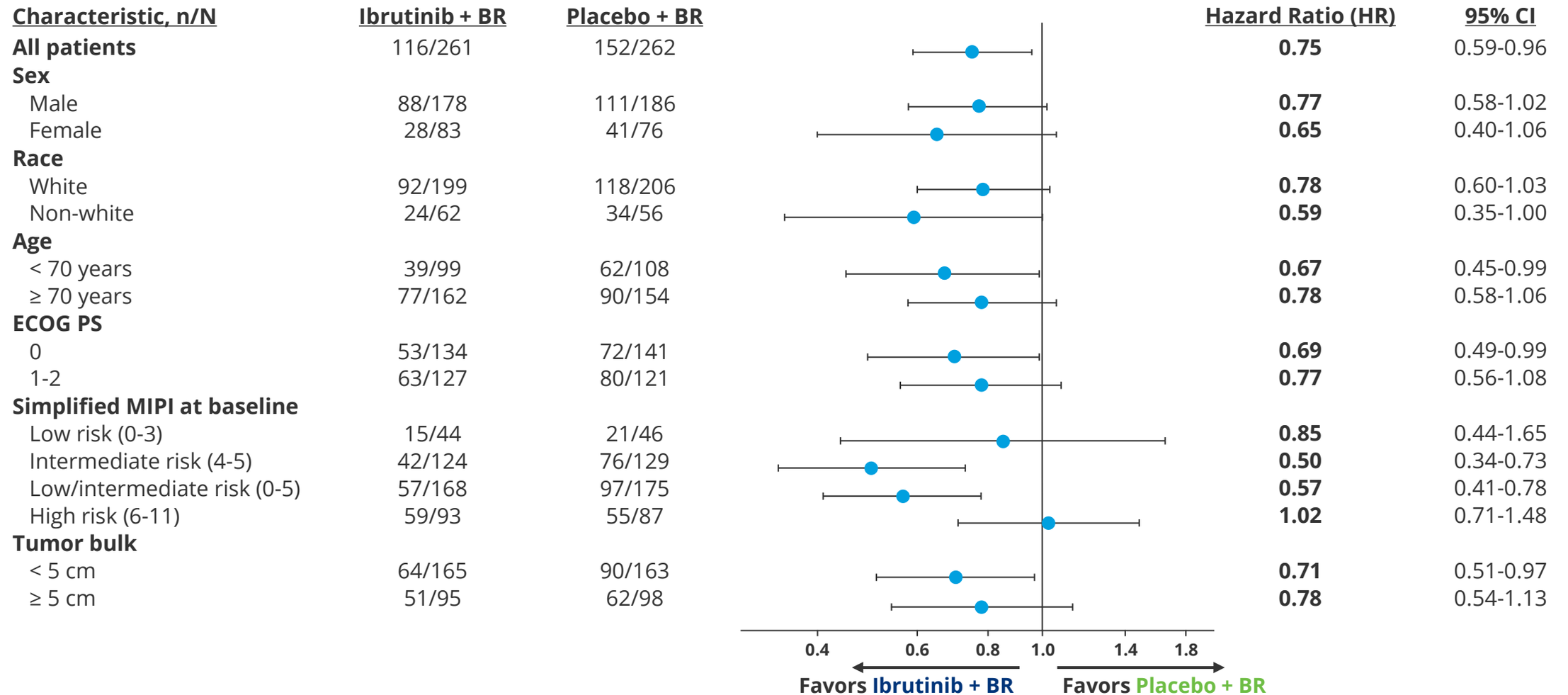
Patients at Risk

Ibrutinib + BR	261	228	207	191	182	167	152	139	130	120	115	106	95	78	39	11	0
Placebo + BR	262	226	199	177	166	158	148	135	119	109	103	98	90	78	41	11	0

CI, confidence interval; HR, hazard ratio; NE, not evaluable.
*Significance boundary for superiority was $p < 0.023$.

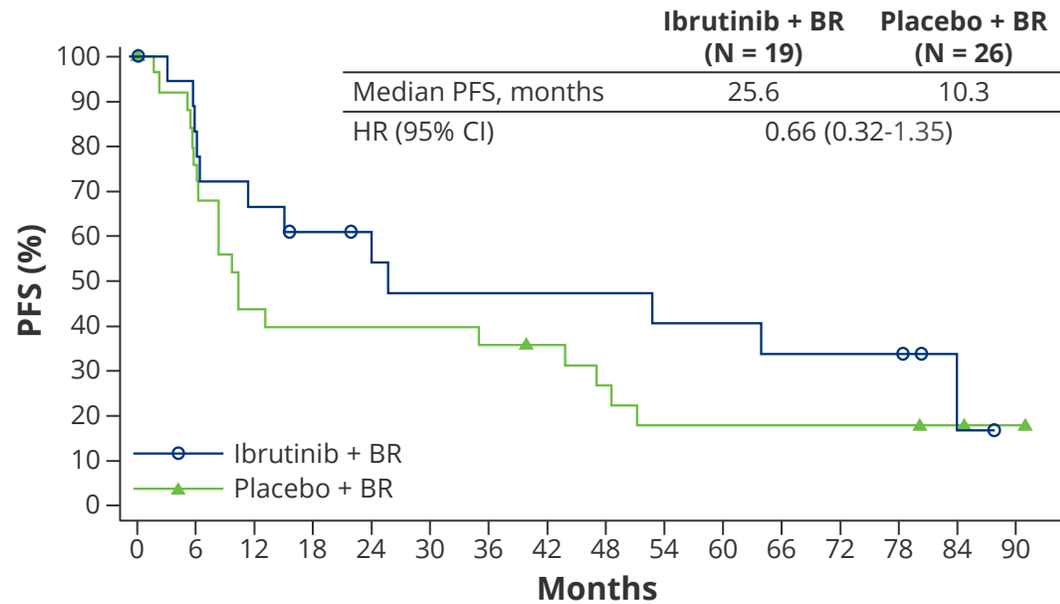


PFS Hazard Ratio in Subgroups



PFS in High-Risk Subgroups

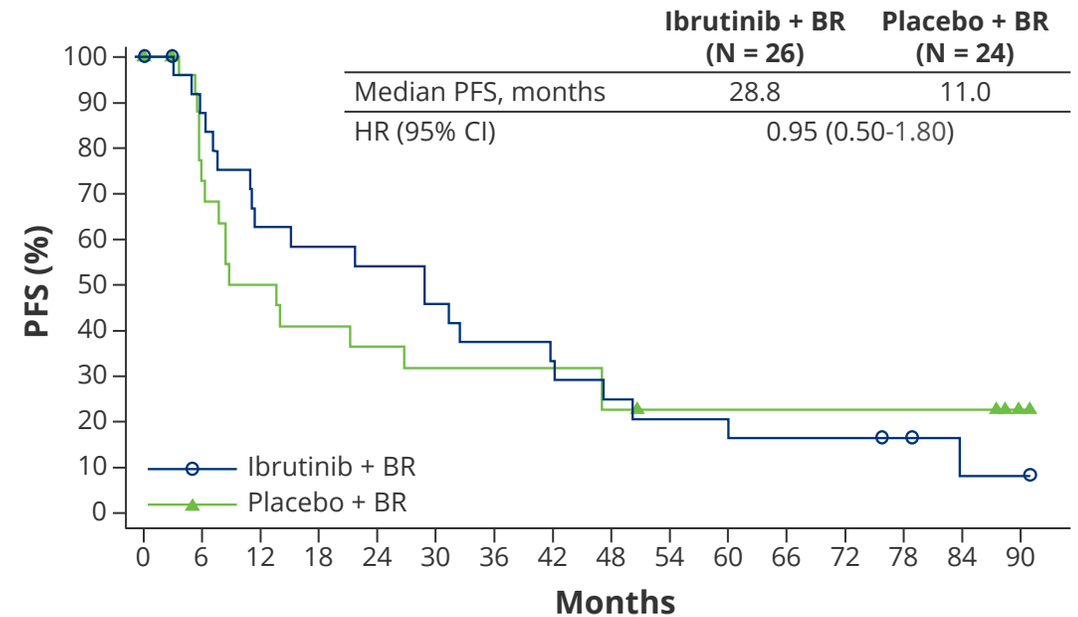
Blastoid/pleomorphic histology



Patients at Risk

Ibrutinib + BR	19	14	12	10	8	7	7	7	7	6	6	5	5	5	1	0
Placebo + BR	26	19	11	10	10	10	9	8	6	4	4	4	4	4	3	1

TP53 mutation present

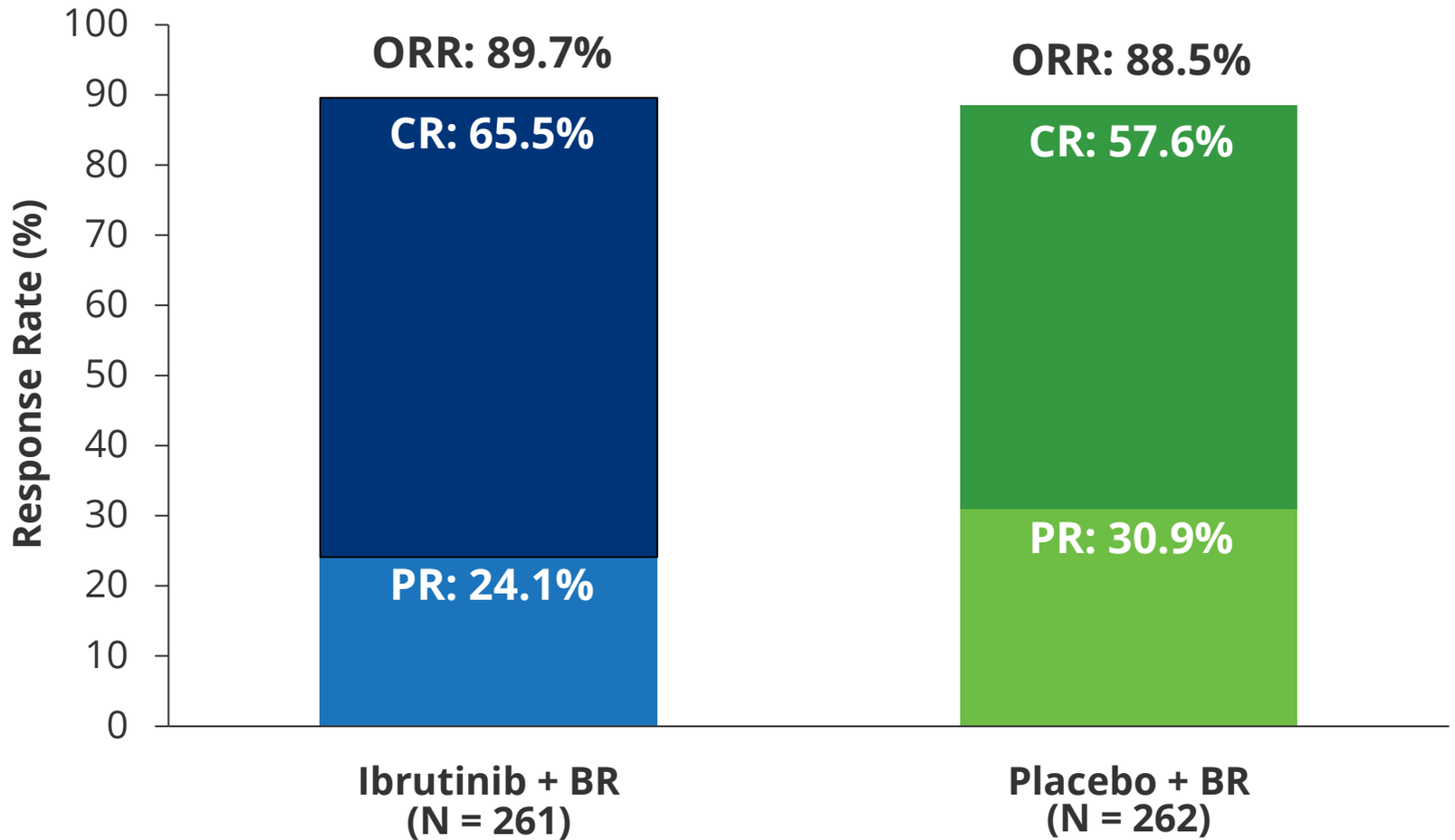


Patients at Risk

Ibrutinib + BR	26	21	15	14	13	11	9	7	6	5	4	4	4	4	3	1	1
Placebo + BR	24	16	11	9	8	7	7	7	5	4	4	4	4	4	4	4	1



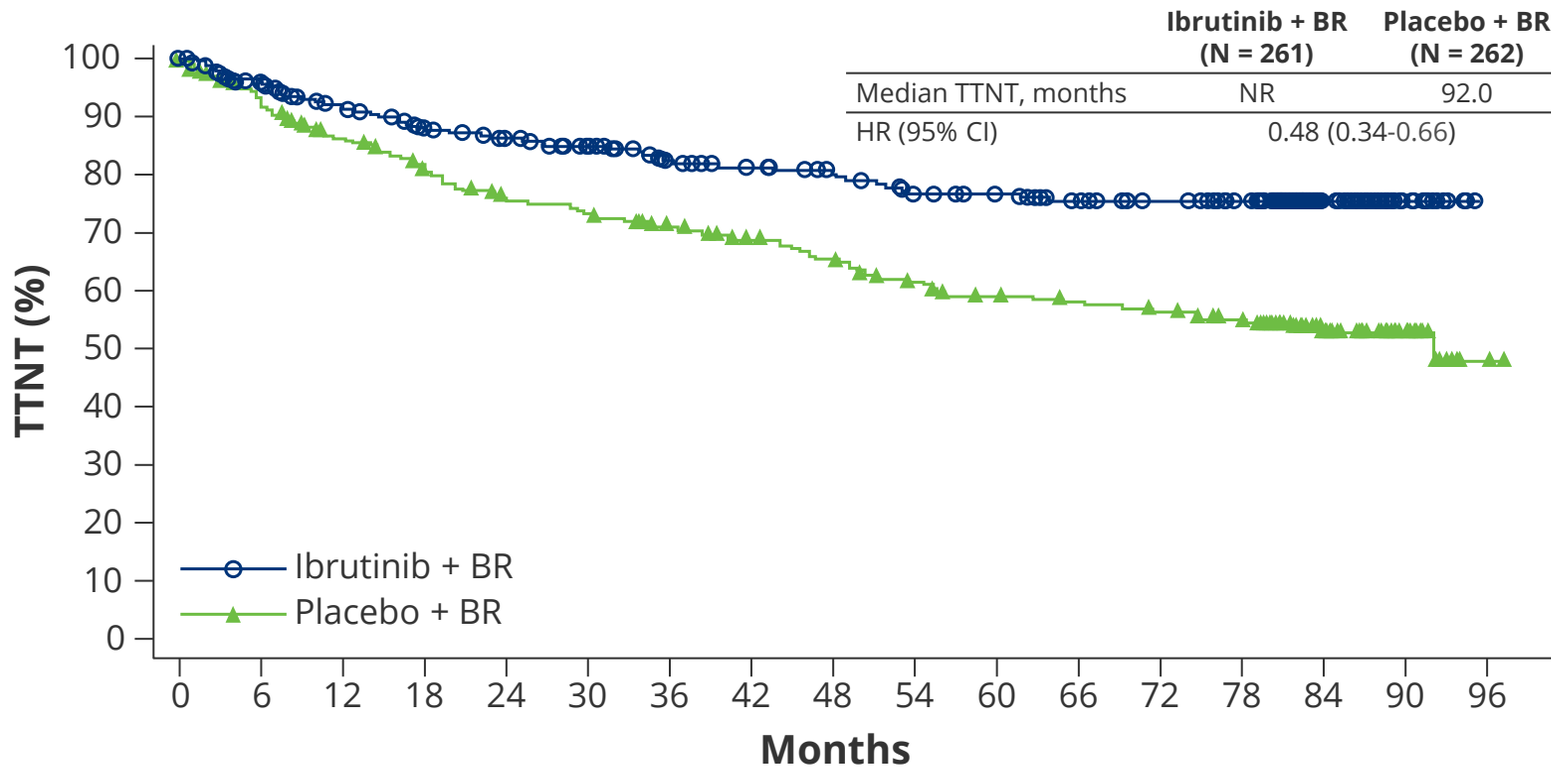
Response Rate



- CR rate was numerically higher in the ibrutinib arm (65.5% vs 57.6%; $p = 0.057$)



Time To Next Treatment



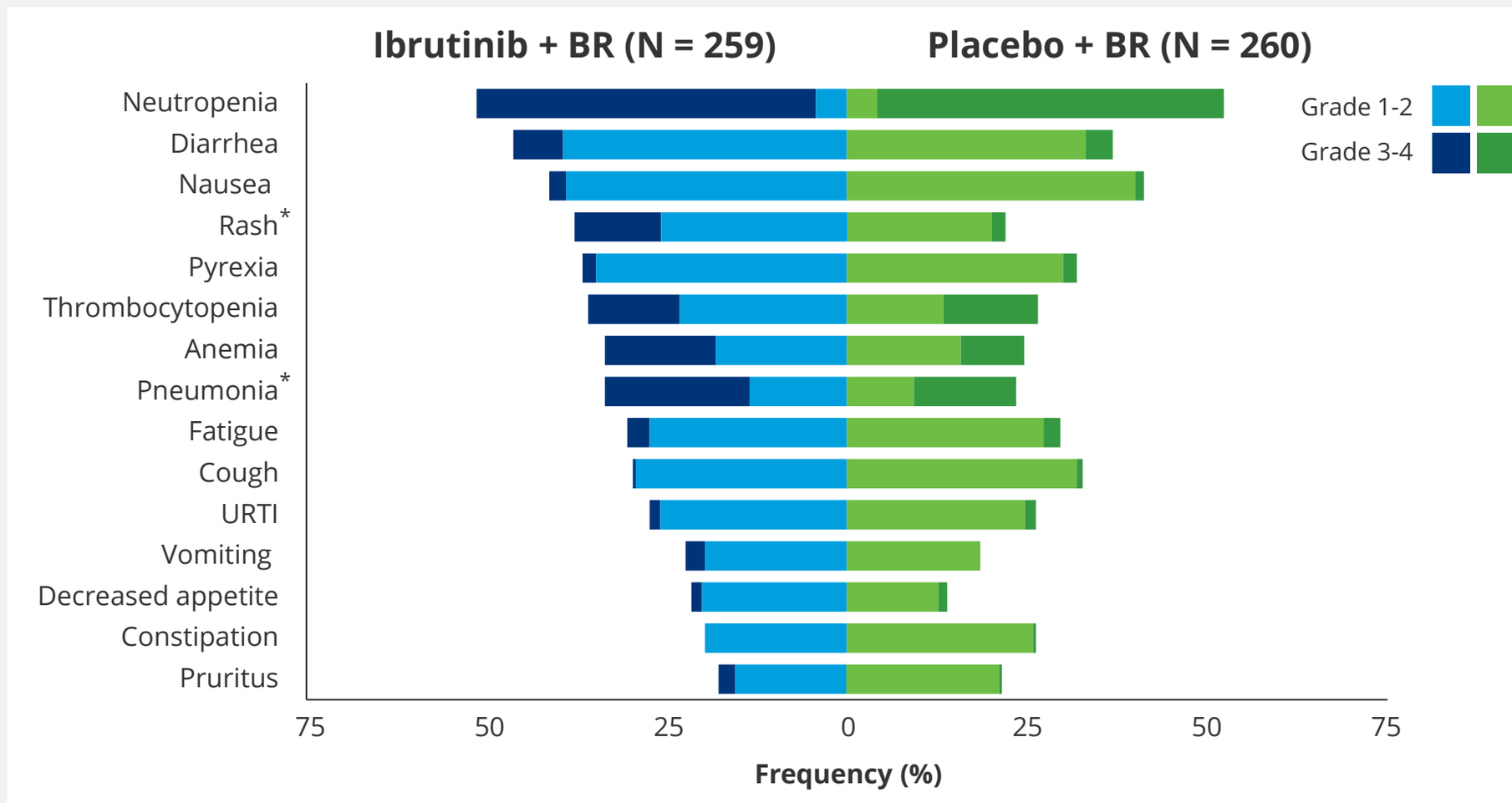
- Subsequent therapy at second-line:
 - Ibrutinib arm: 52/261 (19.9%)
 - BTKi: 6/52 (11.5%)
 - Placebo arm: 106/262 (40.5%)
 - BTKi: 41/106 (38.7%)

Patients at Risk

Ibrutinib + BR	261	231	209	192	184	174	155	147	140	131	126	119	111	102	60	21	0
Placebo + BR	262	231	203	189	171	167	157	146	137	125	117	113	109	101	67	23	2



Common Treatment-Emergent Adverse Events (≥ 20%)



*Difference of ≥ 10% in any grade treatment-emergent adverse event (TEAE).
URTI, upper respiratory tract infection.



TEAEs of Clinical Interest With BTKis

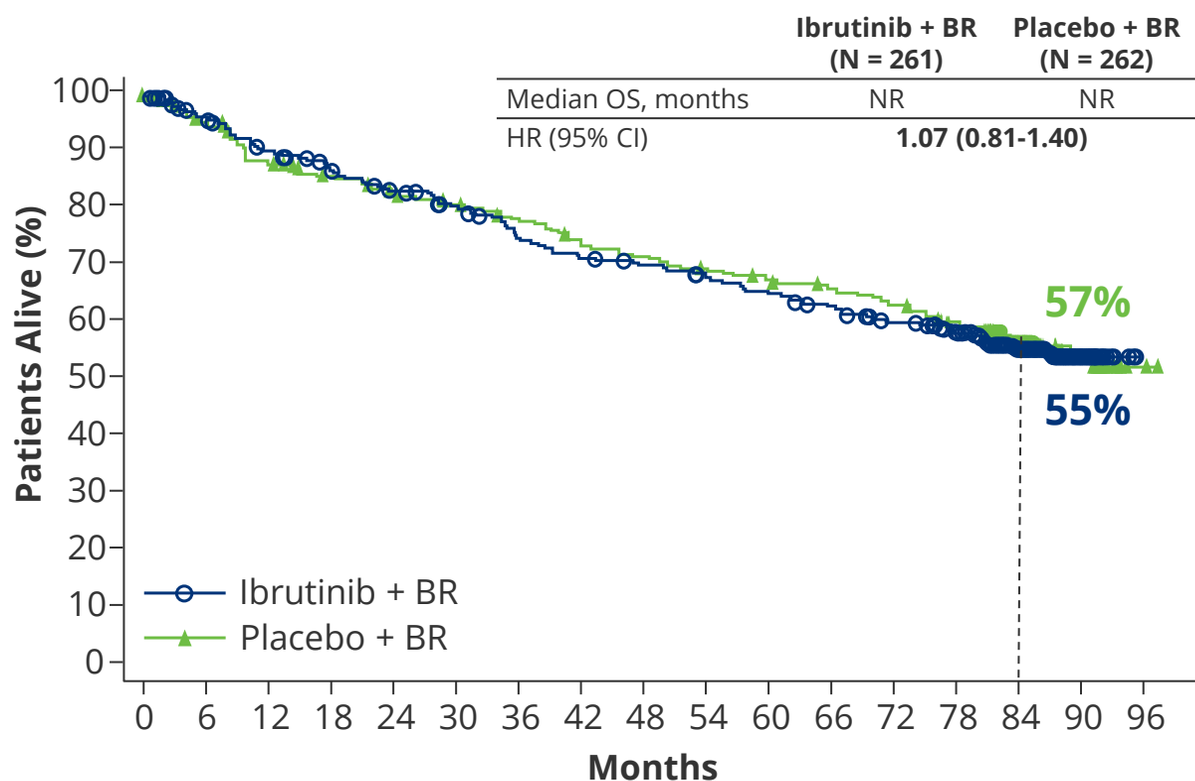
	Ibrutinib + BR (N = 259)		Placebo + BR (N = 260)	
	Any Grade	Grade 3 or 4	Any Grade	Grade 3 or 4
Any bleeding*	42.9%	3.5%	21.5%	1.5%
Major bleeding	5.8%	-	4.2%	-
Atrial fibrillation*	13.9%	3.9%	6.5%	0.8%
Hypertension	13.5%	8.5%	11.2%	5.8%
Arthralgia	17.4%	1.2%	16.9%	0

- These adverse events were generally not treatment limiting
- During the entire study period, second primary malignancies (including skin cancers) occurred in 21% in the ibrutinib arm and 19% in the placebo arm; MDS/AML in 2 and 3 patients, respectively

*Difference of $\geq 5\%$ in any grade TEAE; MDS/AML, myelodysplastic syndromes/acute myeloid leukemia; Any bleeding is based on Haemorrhage Standardized MedDRA Query (SMQ) (excluding laboratory terms). Major bleeding includes any grade 3 or higher bleeding and serious or central nervous system bleeding of any grade.



Overall Survival



Patients at Risk

Ibrutinib + BR	261	239	221	208	197	187	171	163	158	152	145	138	128	118	70	25	0
Placebo + BR	262	244	223	212	203	197	188	177	171	165	159	154	147	137	90	31	2

Cause of death	Ibrutinib + BR (N = 261)	Placebo + BR (N = 262)
Death due to PD and TEAE	58 (22.2%)	70 (26.7%)
Death due to PD	30 (11.5%)	54 (20.6%)
Death due to TEAEs*	28 (10.7%)	16 (6.1%)
Death during post-treatment follow-up excluding PD and TEAEs	46 (17.6%)	37 (14.1%)
Total deaths	104 (39.8%)	107 (40.8%)

- Death due to Covid-19: 3 patients in the ibrutinib arm during the TEAE period and 2 patients in the placebo arm after the TEAE period
- Exploratory analysis of cause-specific survival including only deaths due to PD or TEAEs showed an HR of 0.88

*The most common grade 5 TEAE was infections in the ibrutinib and placebo arms: 9 versus 5 patients. Grade 5 TEAE of cardiac disorders occurred in 3 versus 5 patients, respectively. CI, confidence interval; HR, hazard ratio; NR, not reached; PD, progressive disease; TEAE, treatment-emergent adverse event.



Conclusions

SHINE is the first phase 3 study to show that ibrutinib in combination with chemoimmunotherapy is highly effective in patients with untreated MCL

Median PFS of 6.7 years: a statistically significant and clinically meaningful 2.3-year PFS advantage



Consistent and expected AEs with the known profiles of ibrutinib and BR



A new benchmark for first-line treatment of older patients with MCL or those unsuitable for ASCT



Thanks to all my co-authors

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Acknowledgements

The SHINE study team would like to thank the patients who participated in the study and their families, all investigators and personnel at 183 study sites in 28 countries, and members of the SHINE independent data monitoring committee.

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Argentina

Dorotea Fantl, Maria Flores, Maria Cecilia Foncuberta, Gustavo Jarchum, Mauricio Leonardo Kotliar, Romina Mariano, Miguel Arturo Pavlovsky

Australia

Cecily Forsyth, Pratyush Giri, Anna Johnston, Hock Choong Lai, Joseph McKendrick, James Morton, Andrew Spencer, Judith Trotman

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Marc Andre, Jan Lemmens, Fritz Offner, Sylvia Snauwaert, Eric Van Den Neste, Achiel Van Hoof, Vibeke Vergote, Gregor Verhoef, Ka Lung Wu

Brazil

Wolney Barreto, George Barros, Marcelo Eduardo Zanella Capra, Carlos Chiattonne, Patricia Giacon, Iara Gonçalves, Alexandre Palladino, Juliana Pereira, Guilherme Perini, Eduardo Rego, Rodrigo Santucci Alves da Silva, Adriana Scheliga, Renato Tavares, Luciana Viola

Canada

Tom Kouroukis, Randeep Sangha, John M. Storrington, Richard Van Der Jagt, Diego Villa

China

Weijun Fu, Xiaonan Hong, Jian Hou, Huiqiang Huang, Jie Jin, Xiaoyan Ke, Junmin Li, Ting Liu, Jianhui Qiao, Lugu Qiu, Hanyun Ren, Yuankai Shi, Yuqin Song, Huaqing Wang, Zhao Wang, Huilai Zhang, Daobin Zhou, Jun Zhu

Czech Republic

David Belada, Jiri Mayer, Heidi Mocikova

France

Kamal Bouabdallah, Caroline Dartigeas, Richard Delarue, Thomas Gastinne, Remy Gressin, Corinne Haioun, Olivier Hermine, Steven Le Gouill, Catherine Thieblemont

Germany

Martin Dreyling, Andreas Loew, Corinna Leng, Julia Meissner, Michaela Schwarz, Ernst Späth-Schwalbe, Stephan Stilgenbauer, Stefan Wirths

Greece

Achilles Anagnostopoulos, Meletios Dimopoulos, Panagiotis Panagiotidis, Vasiliki Pappa

Hungary

Zita Borbenyi, Miklos Egyed, Arpad Illes, Zsolt Nagy, Andras Rosta, Arpad Szomor

Ireland

Amjad Hayat, Elisabeth Vandenberghe

Israel

Irit Avivi, Andrei Braester, Yossef Cohen, Neta Goldschmidt, Ronit Gurion, Yair Herishanu, Maya Koren-Michowitz, Itai Levi, Arnon Nagler, Shimrit Ringelstein, Avichai Shimoni, Tamar Tadmor

Italy

Carola Boccomini, Andrés José Maria Ferreri, Ferdinando Frigeri, Gianluca Gaidano, Marco Gobbi, Roberto Massimo Lemoli, Maurizio Martelli, Antonio Pinto, Alessandro Rambaldi, Umberto Vitolo, Pier Luigi Zinzani

Japan

Noriko Fukuhara, Kiyohiko Hatake, Michiko Ichii, Tatsuo Ichinohe, Kenichi Ishizawa, Koji Kato, Dai Maruyama, Yuko Mishima, Hirohisa Nakamae, Michinori Ogura, Hirohiko Shibayama, Masafumi Taniwaki, Yasuhito Terui, Takanori Teshima, Toshiki Uchida

Korea, Republic of

June-won Cheong, Seok-goo Cho, Hyeon Seok Eom, Seok Jin Kim, Cheolwon Suh, Deokhwan Yang, Dok Hyun Yoon

Mexico

David Gomez, Eva Ramirez, Luis Villela

Netherlands

Henriette Berenschot, Eva De Jongh, Jeanette Doorduyn, Marie José Kersten, Hanneke Kluin-Nelemans, Monique Minnema, Marcel Nijland, Gustaaf Van Imhoff, Hendrik Veelken

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Ewa Chmielowska, Janusz Halka, Wojciech Jurczak, Wanda Knopinska-Posluszny, Jan Walewski, Tomasz Wrobel

Puerto Rico

Fernando Cabanillas

Russian Federation

Irina Bulavina, Oleg Gladkov, Kamil Kaplanov, Tatiana Klitochenko, Nuriyet Khuazheva, Georgii Manikhas, Alexander Myasnikov, Eugeniy Osmanov, Tatiana Pospelova, Alexander Pristupa, Andrey Proydakov, Olga Samoilova, Olga Serdyuk, Gayane Tumyan, Sergey Voloshin

Slovakia

Juraj Chudej, Andrea Cipkova, Stanislav Palasthy, Andrej Vranovsky, Alexander Wild

Spain

Natalia Alonso, Reyes Arranz, Mariana Bastos, Dolores Caballero, Jorge Gayoso, Armando Lopez Guillermo, José-Ángel Hernández-Rivas, Joan Bargay Lleonart, Concepcion Nicolas, Albert Oriol Rocafiguera

Sweden

Stefanie Baumgartner-Wennerholm, Mats Jerkeman, Claes Karlsson, Ingemar Lagerlöf, Anna Laurell, Karin Papworth

Taiwan

Tsai-yun Chen, Yeu-chin Chen, Bor-sheng Ko, Ching-yuan Kuo, Hsuan-yu Lin, Chun-yu Liu, Po-nan Wang, Su-peng Yeh

Turkey

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United Kingdom

Rebecca Auer, Ian Chau, Martin Dyer, Peter Johnson, Rod Johnson, David Lewis, Pam McKay, Sylvia Montoto, Andrew Pettitt, Chris Pocock, John Radford, Simon Rule, Simon Wagner, Moya Young

United States of America

Ranjana Advani, Ammar Alzoubi, Jennifer Amengual, Bipinkumar Amin, David Andorsky, Anne H. Angevine, Anne Beaven, Maurice Berkowitz, Vipul Bhandari, Lillian Burke, January Castro, Neil Cohen, Kevin David, Christopher Di Simone, Mathew Fero, Roger Fleischman, Ian Flinn, Lawrence Garbo, Andre Goy, Paul A. Hamlin, John Hayslip, Iris Isufi, Mark Kaminski, Aziz Khan, Ali Khojasteh, Leonard Klein, Mouhammed Jameel Kyasa, Brian Link, Delong Liu, Elizabeth McGuire, Matthew McKinney, Madhu Midathada, Emiliano Mugnaini, Ndegwa Njuguna, Gregg Olsen, Kenneth Pennington, Daniel Persky, Adam Petrich, Fahd Quddus, Radhakrishnan Ramchandren, Ruben Reyes, John Reynolds, Jorge Romaguera, Peter Rosen, Lori Rosenstein, Stephen Schuster, Spencer Shao, Jeff Porter Sharman, Gary Spitzer, Julian Sprague, Stephen Spurgeon, Don Stevens, Patrick Stiff, Michael Luhua Wang, Donald Wender, Abdulraheem Yacoub, Jay Yang, Alexander Zweibach

Oral presentation previously presented at the ASCO 2022 Annual Meeting, June 3-7, 2022, Chicago, IL & Online and the EHA2022 Congress, June 9-12, 2022, Vienna, Austria. Also published in NEJM (www.nejm.org/doi/full/10.1056/NEJMoa2201817)

Medical writing and editorial assistance were provided by Ward A. Pedersen, PhD, of Parexel, funded by Janssen Research & Development.

