


Chemotherapy-Free Management of Follicular and Marginal Zone Lymphoma

Thomas A Ollila^{1,2}

Adam J Olszewski ^{1,2}

¹Department of Medicine, Alpert Medical School of Brown University, Providence, RI, USA; ²Division of Hematology-Oncology, Rhode Island Hospital, Providence, RI, USA

Abstract: Many patients with follicular (FL) or marginal zone lymphoma (MZL) are not eligible to receive immunochemotherapy due to advanced age or comorbidities. Recent innovations in the treatment of these indolent lymphomas provide options for multiple lines of chemotherapy-free management. More research is needed to determine which older patients are best served by a chemotherapy-free approach in the context of geriatric vulnerabilities. In the first line, regardless of disease burden, rituximab monotherapy can provide high rates of disease control with minimal toxicity, while judicious use of brief maintenance extends the duration of response. Radioimmunotherapy using ibritumomab tiuxetan is an effective and safe post-rituximab consolidation for older patients who have <25% bone marrow involvement. The combination of rituximab and lenalidomide, although “chemotherapy-free”, does not improve tolerability over immunochemotherapy. However, studies support lower doses and shorter duration of lenalidomide exposure as a means to improve safety without materially compromising efficacy for older individuals. Extranodal MZL can often be effectively controlled with low-dose radiation therapy, and splenic MZL has excellent outcomes with rituximab monotherapy. For many patients with relapsed FL/MZL, simple retreatment with anti-CD20 antibodies will prove sufficient. Other currently available options for relapsed/refractory disease include ibritumomab tiuxetan, lenalidomide with rituximab, umbralisib as a potentially less toxic PI3K inhibitor, ibrutinib (for MZL), and tazemetostat (for FL, especially with *EZH2* mutation). Emerging data with novel forms of immunotherapy (antibody-drug conjugates like polatuzumab vedotin or loncastuximab tesirine; T-cell-engaging bispecific antibodies like mosunetuzumab or epcoritamab; and chimeric antigen receptor CAR T-cells like axicabtagene ciloleucel) suggest that immune-directed approaches can produce very high and potentially durable responses in FL/MZL with limited toxicities, further obviating the need for chemotherapy.

Keywords: follicular lymphoma, marginal zone lymphoma, geriatric oncology, lenalidomide, ibritumomab tiuxetan, bispecific antibodies

Introduction

Indolent B-cell lymphomas (iBCL) are a group of slow-growing mature B-cell lymphomas that often affect older patients. In the United States (US), median age at diagnosis ranges from 65 for follicular lymphoma (FL), 67 for marginal zone lymphoma (MZL), 70 for chronic lymphocytic leukemia/small lymphocytic lymphoma (CLL/SLL), to 72 for lymphoplasmacytic lymphoma/Waldenström macroglobulinemia (LPL/WM).¹ Many older patients with iBCL will have comorbid conditions and functional impairments that make treatment with cytotoxic chemotherapy difficult.² Furthermore, advanced iBCL are not considered curable by standard therapy, yet average life expectancy is measured in years to decades and

Correspondence: Adam J Olszewski
Department of Medicine, Alpert Medical School of Brown University, Rhode Island Hospital, 593 Eddy St., Providence, RI, 02903, USA
Tel +1 844-222-2881
Fax +1 401-444-8918
Email adam_olszewski@brown.edu

many patients will receive several lines of treatment. These factors highlight the importance of treatment choices that balance efficacy against toxicity considering patient's expected lifetime.

The 3 most common types of iBCL are FL (grade 1 to 3a), MZL, and SLL. Grade 3b FL is considered more aggressive and is managed using strategies appropriate for diffuse large B-cell lymphoma (DLBCL). MZL is further subclassified into nodal MZL (NMZL), splenic MZL (SMZL), and extranodal MZL (EMZL) of the mucosa-associated lymphoid tissue (MALT lymphoma).³ These subtypes have specific clinical features and management approaches. LPL/WM is uniquely characterized by complications of monoclonal IgM paraproteinemia, and its treatment follows specific patterns.⁴ In this review, we do not discuss CLL/SLL and LPL/WM, which follow disease-specific treatment pathways.

For the past two decades, the management of FL and MZL has relied on a few general principles.^{5,6} Early-stage disease (stage 1/2) could be treated with potentially curative intent using radiation therapy (or, in certain types of MZL, excision or antibiotic therapy).^{7–10} Asymptomatic patients with advanced stage disease would be typically observed without treatment or offered single-agent rituximab.^{11,12} Patients with a higher burden of disease would be mostly offered rituximab-containing immunochemotherapy. One current standard first-line immunochemotherapy approach is based on the Phase 3 German Study group indolent Lymphomas (StiL) trial, which showed improved progression-free survival (PFS) and fewer acute toxicities with rituximab plus bendamustine (BR) compared with the historical standard of R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone).¹³ However, even BR is associated with 29% risk of grade 3/4 neutropenia and 37% risk of infections. Furthermore, it leads to a prolonged T-cell depletion and potential opportunistic infections, including zoster, pneumonia, or CMV reactivations, particularly among older patients.^{14–16} Many older patients are thus understandably weary of such risks and prefer to avoid chemotherapy.

Recent advances allow older and unfit patients to expect reasonable efficacy with more tolerable, targeted therapies, which, if adequately sequenced, may obviate the need for any chemotherapy within their lifetime. In this review, we aim to present both established and novel chemotherapy-free approaches to systemic treatment for

FL and MZL, which are particularly (but not exclusively) relevant for older patients.

Defining Candidates for Chemotherapy-Free Approach

When treating an older patient with iBCL, clinicians face the complex question of whether the potential benefits of cytotoxic chemotherapy might be outweighed by risks. Geriatric assessment (GA) can help clarify this balance, although its use is less frequent in hematologic malignancies than in solid tumor oncology.^{17–19} Management of iBCL requires nuance to help decide if, when, and which therapy should be started, in the context of multiple options of variable intensity. The watch-and-wait approach can be applied to patients with asymptomatic and low-volume disease.¹¹ The Groupe d'Etude des Lymphomes Folliculaires (GELF) or British National Lymphoma Investigation (BNLI) criteria provide objective definitions of high-burden FL.^{20,21} Prognostic scores like the FL International Prognostic Index (FLIPI) or MALT IPI may predict expected survival, but do not define an indication for treatment or a benefit from specific regimens.^{22–24} A dedicated GA can help guide the decision-making, providing insight into functional and nutritional status, polypharmacy, as well as psychosocial factors. GA has been used in diffuse large B-cell lymphoma (DLBCL) for selection of treatment intensity, but data in iBCL are sparse.^{25,26} Observational studies have demonstrated that GA predicts shorter survival, treatment discontinuation, and the risk of hospitalization beyond simple "performance status" assessments, mostly in the context of cytotoxic chemotherapy.²⁷ Clinical trials of chemotherapy-free approaches in indolent lymphomas have so far not used GA for patient selection. However, a recent Phase 2 study has demonstrated that patients age 65–80 who scored "fit" on the simplified comprehensive GA could be safely treated with a short course of immunochemotherapy (rituximab, bendamustine, plus mitoxantrone), achieving 78% rate of complete response (CR) and a 3-year PFS of 67%.²⁸ Only 4 of 72 participants discontinued chemotherapy for toxicity.

When comprehensive GA is not available, hematologists can assess patients' fitness using simple tools like the Timed Up and Go (TUG) test, which involves brisk walking back and forth for 3 meters from a sitting position.¹⁸ Needing >30 seconds to complete the TUG test correlates with significant frailty. In a prospective observational study of patients with hematologic malignancies (29% various lymphomas), gait

Table 1 Chemotherapy-Free Options for First-Line Treatment in FL or MZL

Treatment	Disease	N	Age, Median (Range)	Gr. 3/4 AE	ORR	CR/CRu	Median PFS (y)
Rituximab							
IR ^{32,33}	FL	50	52 (32–75)	4%	73%	26%	2.0
IR ¹¹	FL	84	60 (33–86)	6%	77%	47%	>4 ^a
IR ^{34,35}	FL/MZL	321	57 (29–82)	7% ^b	74% ^b	41% ^b	1.45 ^c
MR ¹¹	FL	192	60 (27–87)	11%	88%	69%	>5 ^a
MR ³⁷	FL	270	55 (25–82)	9%	63%	16%	7.4
IR ± MR ³⁸	FL	289	59 (25–86)	5%	71%	12%	~3 ^a IR, NR MR
IR ± MR ³⁹	MZL/SLL	71 MZL	66 (30–86)	6%	52% (MZL)	13% (MZL)	1.4 IR, 4.8 MR ^c
IR ⁶⁹	EMZL	138	63 (27–81)	10%	78%	56%	6.9
IR ± MR ⁷⁵	SMZL	108	65 (41–91)	^a	92%	65%	> 10
Rituximab + lenalidomide							
R2 ⁵²	FL	413	59 (30–89)	65%	61%	48%	77% at 3 y
R2 ⁵⁷	FL	77	61 (26–80)	56%	81%	36%	5.0
R2 ⁵⁶	MZL	30	58 (36–77)	≥ 33% ^a	93%	70%	5.0
Ibritumomab tiuxetan							
⁹⁰ YIT ⁶⁰	FL	50	60 (37–81)	≥ 30% ^a	94%	86%	63% at 3 y
⁹⁰ YIT ⁶¹	FL, MZL	31	57 (28–87)	≥ 61% ^a	100%	97%	~75% at 3 y ^a
⁹⁰ YIT ⁶²	FL	59	66 (51–83)	48%	87%	56%	2.2
⁹⁰ YIT ⁶⁶	FL	74	61 (28–80)	56%	96%	69%	3.4
⁹⁰ YIT ⁶⁷	MZL	16	62 (37–84)	50%	88%	56%	4.0

Notes: ^aNot explicitly reported; ^brituximab monotherapy arm; as reported in the MLI6865 trial;³⁵ ^ctime to treatment failure.

Abbreviations: AE, adverse event; CR/CRu, complete response/complete response unconfirmed; DOR, duration of response; EFS, event-free survival; EMZL, extranodal marginal zone lymphoma; IR, induction rituximab; MR, maintenance rituximab; NR, not reached; ORR, overall response rate; PFS, progression-free survival; R2, rituximab and lenalidomide; TTF, time to treatment failure; y, years; ⁹⁰YIT, ibritumomab tiuxetan.

speed measured over a 4-meter walk strongly correlated with overall survival, unplanned hospitalizations, and emergency room visits—but did not appear to influence treatment intensity recommendation.^{29,30} Although clinical data on indolent lymphomas are lacking, in the CLL9 clinical trial of cytotoxic chemotherapy, the TUG test was a better predictor of survival than the assessment of comorbidities and of the instrumental activities of daily living.³¹ We recommend that clinicians formally assess their older patients with iBCL for functional capacity and that they consider chemotherapy-free regimens for frail individuals. On the one hand, age alone should not disqualify a fit older patient from receipt of chemotherapy, but on the other—patients with functional impairments and comorbidities may be the group benefitting most from treatment options that avoid cytotoxic chemotherapy.

First-Line Therapy for FL and MZL

Evidence for the use of chemotherapy-free treatments in iBCL is largely based on experience from FL, but some MZL-specific studies support their use for different MZL

subtypes (Table 1). The principal options include single-agent rituximab, the combination of rituximab with lenalidomide, and radioimmunotherapy (Figure 1).

Anti-CD20 Antibodies: Selective Use of Maintenance

Rituximab is a chimeric monoclonal anti-CD20 antibody with over 20 years of cumulative experience in iBCL. First-line rituximab monotherapy in low-burden FL results in an overall response rate (ORR) of 73%, a rate of complete response (CR) or complete response unconfirmed (CRu) of 26% at the end of therapy which deepens to 52% over time, median PFS of 24 months, and median duration of response (DOR) of 29 months.^{32,33} The disease control is also excellent for symptomatic patients: in a long-term follow-up of 2 Nordic Lymphoma Group trials of rituximab (± interferon alpha) in symptomatic iBCL (84% FL), 36% of patients never needed chemotherapy, and OS at 10 years was 75%—similar to trials that used first-line immunochemotherapy.^{34,35} One international randomized study compared watchful waiting with rituximab

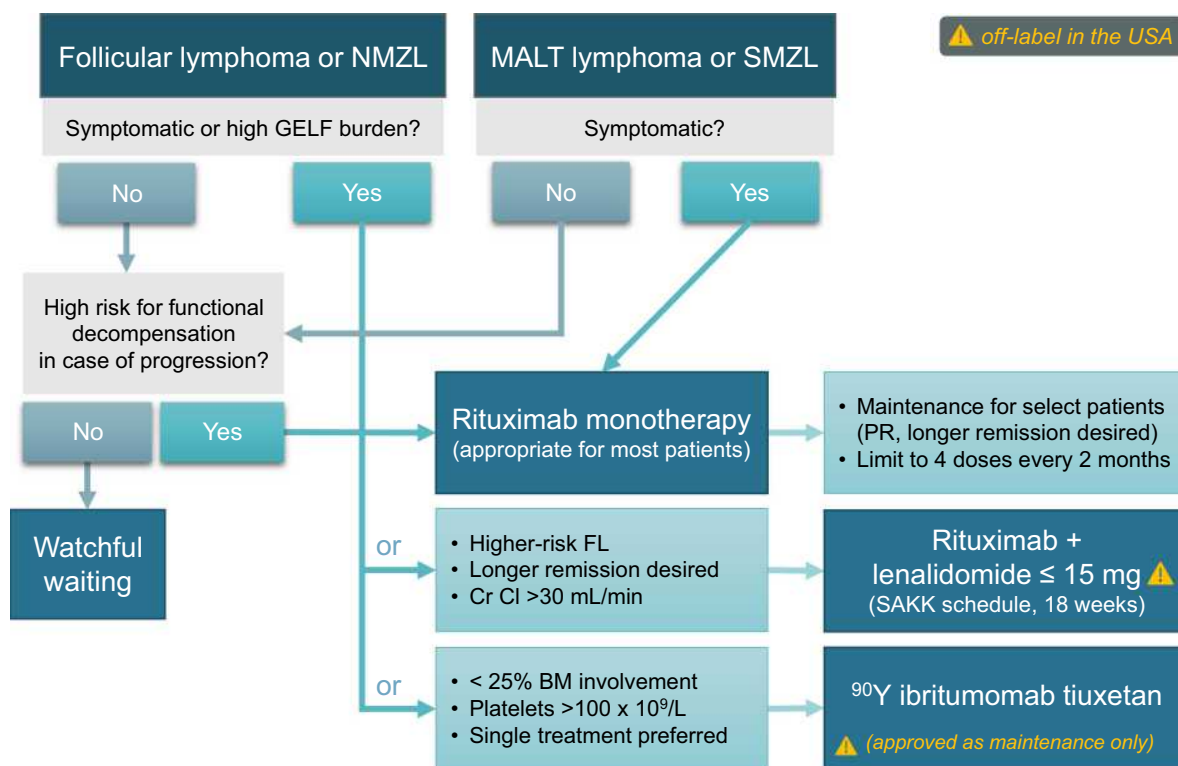


Figure 1 Chemotherapy-free options for first-line systemic therapy of FL or MZL (note that some options are off-label in the USA).

Abbreviations: BM, bone marrow; CrCl, creatinine clearance; FL, follicular lymphoma; GELF, Groupe d'Etude des Lymphomes Folliculaires; MALT, mucosa-associated lymphoid tissue; NMZL, nodal marginal zone lymphoma; PR, partial response; SAKK, Swiss Group for Clinical Cancer Research; SMZL, splenic marginal zone lymphoma.

± 2-year maintenance in low-burden FL.¹¹ Following a single 4-week course of rituximab, 78% of participants did not require any therapy for over 3 years; this proportion was even higher (88%) if maintenance was applied. The overall quality of life was similar among patients undergoing watchful waiting versus rituximab therapy, except for improved measures of mental adjustment to cancer and illness coping with active therapy.¹¹ In a phase 2 trial by the Swiss Group for Clinical Cancer Research (SAKK) which enrolled patients regardless of disease burden or prior chemotherapy, a brief (4 doses every 2 months) rituximab maintenance also extended median event-free survival (EFS) to 24 months compared with 13 months with rituximab induction alone.³⁶ However, longer maintenance (up to 5 years) did not provide additional benefit.³⁷ Similarly, the E4402 trial randomized patients with FL or other iBCL (including MZL) to maintenance rituximab or simple retreatment in case of progression, showing no significant difference in median time to treatment failure (4.3 versus 3.9 years for FL, $P=0.54$) or quality of life.^{38–40} Therefore, the potential benefits of maintenance rituximab should be balanced against the burden of extended therapy and associated

infectious toxicity, particularly for patients with other risk factors for infections.

The subcutaneous formulation of rituximab with hyaluronidase may provide similar efficacy with increased convenience, although it is not currently approved in the first-line setting as monotherapy.⁴¹ Given the established efficacy of the formulation, it can be considered after at least 1 intravenous dose of rituximab is given without a severe infusion reaction. Recent data also support the use of rituximab biosimilars for first-line therapy.⁴² Of note, another anti-CD20 antibody ofatumumab failed to show improvement in outcomes over rituximab.^{43,44}

Obinutuzumab is a type II glycoengineered anti-CD20 IgG monoclonal antibody with increased direct cytotoxic effect as well as antibody-dependent cellular cytotoxicity.⁴⁵ It has shown efficacy for iBCL in both relapsed/refractory and first-line settings, extends PFS when combined with chemotherapy instead of rituximab, and has gained FDA approval in combination with chemotherapy for treatment of FL.^{46–49} However, when used as monotherapy in the phase 2 randomized GAUSS trial in FL, obinutuzumab increased ORR without any PFS or toxicity advantage over rituximab.⁴⁷ Therefore, we recommend rituximab rather

than obinutuzumab for first-line monotherapy in iBCL, with a selective use of maintenance for responding patients who value a longer remission over potential retreatment.

Lenalidomide and Rituximab: Lower Dose, Shorter Duration

Lenalidomide in combination with rituximab (R2) has shown high activity in FL and MZL and is currently FDA-approved for these histologies after at least 1 line of therapy. In previously untreated patients, phase 2 studies showed ORR exceeding 90% and CR rates exceeding 70%.^{50,51} The subsequent phase 3 RELEVANCE trial compared R2 with chemoimmunotherapy (RCHOP in 72%) in untreated FL.⁵² Lenalidomide was dosed at 20 mg daily on days 1–21 for 6 four-week cycles, and then decreased to 10 mg for additional 12 months in patients with CR/CRu, for a total of 18 months of therapy. Rituximab was given weekly during cycle 1, monthly for cycles 2–6, and every 8 weeks for additional 12 cycles. However, the overall toxicity profile was not lower than with immunochemotherapy. In fact, more patients receiving R2 had dose reductions, interruptions, or early discontinuations, and 67% experienced grade 3/4 adverse events which included neutropenia in 32% and rash/cutaneous reactions in 7%. Less severe, but important (considering prolonged exposure) toxicities included anemia (66%), diarrhea (37%), fatigue (23%), nausea (20%), abdominal pain (15%) and myalgias (14%). Although late or longitudinal toxicities are insufficiently evaluated in clinical trials, experience with lenalidomide in FL from the Alliance 50401 trial suggests that fatigue in lenalidomide decreases over time, but neutropenia is cumulative and steady.⁵³ No excess in secondary cancers was reported in the RELEVANCE trial.⁵² With no significant advantage over chemotherapy in PFS (HR, 1.10; 95% CI 0.85–1.43), other efficacy or toxicity endpoints, and the need for 1.5 years of continuous therapy, the R2 regimen, while technically “chemotherapy-free,” cannot be considered an improvement for older/unfit patients with FL. It also could not be safely combined with a third targeted agent like ibrutinib or idelalisib.^{54,55} In MZL, R2 retains a high efficacy with reported ORR of 93%, CR of 70%, and median PFS of 60 months, but the toxicity profile is similar to FL.⁵⁶

Importantly, the immunomodulatory effects of lenalidomide can be achieved with lower doses of the drug. The SAKK 35/10 phase 2 trial randomized 154 previously

untreated FL patients to two 4-week courses of rituximab given 3 months apart, with or without lenalidomide 15 mg administered continuously for 18 weeks.⁵⁷ Lenalidomide was started 2 weeks before the first infusion of rituximab and further dose-reduced to 5 mg for creatinine clearance <60 mL/min. The ORR to this R2 version was 81% and the CR/CRu was 36% (61% by independent review), with median PFS of 5 years—all better than for rituximab alone. Grade ≥ 3 toxicity was higher with lenalidomide (56% versus 22%), but grade 3/4 neutropenia (23%) appeared less frequent than in the RELEVANCE trial and no other grade 3/4 toxicity occurred in more than 5%. Common lower-grade adverse events included fatigue (52%), infections (30%), rash (27%), diarrhea (25%), cough (25%), and nausea/vomiting (22%). These results suggest that addition of lenalidomide to rituximab can provide a benefit for older patients who need higher ORR or longer duration of remission, but lenalidomide can be administered at lower doses and for a short duration. The immunomodulatory effects of lenalidomide among older patients need to be investigated in further research considering immune senescence. It is important to note that lenalidomide increases the risk of thromboembolism and thromboprophylaxis should be considered.

Radioimmunoconjugates

Although maintenance rituximab can extend PFS after the initial induction, an alternative, albeit underused, approach involves consolidation using the ⁹⁰Y radioimmunoconjugate ibritumomab tiuxetan (⁹⁰YIT). ⁹⁰YIT was initially studied after cytotoxic chemotherapy,^{58,59} but it has also been applied as a single-agent for early- and advanced-stage iBCL.^{60–62} ⁹⁰YIT has the potential advantage of delivery as a single course completed within 10 days, which includes the 2 rituximab doses required 7–9 days before and on the day of ⁹⁰YIT infusion. Because of the potential for toxicity resulting from bone marrow irradiation, ⁹⁰YIT can only be applied to patients with a platelet count >100 x10⁹/L and less than 25% lymphomatous infiltration of the bone marrow. Concerns about secondary myeloid malignancy often relegate the use of ⁹⁰YIT to subsequent lines of therapy, but the risk is actually not increased when ⁹⁰YIT is given before any exposure to cytotoxic chemotherapy.^{60,63–65} In phase 2 studies, ORR in FL was 87–94%, CR rate was 56–86%, PFS at 3 years was 58–63%, and overall survival was >90%.^{60,62,66} Grade 3/4 non-hematologic toxicities were exceptionally rare,

although neutropenia or thrombocytopenia were noted in up to 30%, with nadir at ~35 days and universal resolution after 14 weeks from treatment. ^{90}YIT is equally effective in MZL, but it is only approved for relapsed/refractory iBCL or as consolidation in FL after first-line chemotherapy.⁶⁷

^{90}YIT provides an attractive single-course therapy for iBCL for older patients given its efficacy, relatively mild non-hematologic toxicity and low burden of treatment visits, but it requires coordination between the hematologist and the nuclear medicine department and is not universally available. In addition, the requirement for baseline bone marrow assessment and for monitoring of the extended myelosuppression may pose barriers for some older individuals. ^{90}YIT has not been studied in SMZL, which typically extensively involves the bone marrow, and it should be used with caution in this disease.

Special Cases: SMZL and MALT Lymphoma

While immunochemotherapy is typically applied to FL and NMZL, rituximab monotherapy is often sufficient for advanced MALT lymphoma or SMZL regardless of age, with no detriment to survival and with less toxicity when chemotherapy is omitted.^{2,68–71} In the phase 3 study (IELSG-19) comparing first-line systemic chlorambucil, rituximab, or both drugs in MALT lymphoma, single-agent rituximab has shown ORR of 78%, CR rate of 56%, and median event-free survival (EFS) of 5.6 years.⁶⁹ An important consideration for older and unfit patients is the high radiosensitivity of MALT lymphoma, which can often be effectively controlled with minimal doses (4 Grays) of radiation therapy delivered to single or multiple sites of disease.⁷²

Rituximab is also an excellent first-line option for SMZL obviating the use of splenectomy for most patients.^{71,73,74} In fact, observational data suggest no benefit of the addition of chemotherapy to the monoclonal antibody in this disease for older individuals.⁷⁰ The largest phase 2 trial used a 6-week rituximab induction followed by 1–2 years of maintenance. It reported ORR of 92%, CR/CRu of 65%, and 5-year PFS of 71%.⁷⁵ Patients with SMZL should always be screened for the presence of hepatitis C, as regression of the lymphoma has been noted after treatment for this infection.^{76,77} For patients unsuitable for any systemic therapy or splenectomy, radiation is a seldom used but efficacious option.⁷⁸

Relapsed/Refractory Disease

Among patients who experience relapsed or refractory FL/MZL after initial therapy, the extent of symptoms and disease burden, duration of first response (< or \geq 2 years), as well as preference for intravenous or orally administered therapy will determine further management (Figure 2).^{79,80} Clinical trials are always a consideration in this setting and should be offered to eligible older and less fit patients. Many patients with iBCL experience asymptomatic or slow progression which might not require immediate therapy—though, as in the first-line setting, the risk of decompensation during watchful waiting should be considered in the context of comorbidities, psychosocial support, tumor location and bulk. Rapidly progressive recurrence should prompt evaluation of a histologic transformation. Clinicians should also remember that low-intensity chemotherapy (for example, chlorambucil or judiciously dosed BR) can control FL or MZL for many patients with toxicity that is no worse, and sometimes better, than some “chemotherapy-free” approaches.^{21,69,81–83}

Retreatment with Rituximab

Patients initially treated with single-agent rituximab and not on maintenance can be often simply retreated upon progression according to the RESORT trial paradigm.³⁸ In a phase 2 trial, retreatment with rituximab (at median 14.5 months from the previous course) resulted in ORR of 40%, including 11% CR, and an estimated median time to progression of 18 months.⁸⁴ Using an alternative antibody (ofatumumab or obinutuzumab) does not appear to meaningfully improve outcomes (Table 2).^{43,47} However, for patients who experience a rituximab-refractory FL/MZL or a symptomatic relapse within 2 years of prior therapy, we favor an alternative approach when life expectancy exceeds 2 more years because a second remission with rituximab alone is likely to be brief.

Radioimmunoconjugates

The use of ^{90}YIT in relapsed iBCL is FDA-approved and supported by phase 2 and 3 trials.^{85–87} Among patients with rituximab-refractory FL (defined as no response or progression within 6 months), ORR to ^{90}YIT is 74%, CR rate is 15%, and median time to progression is 7 months.⁸⁸ In a subsequent phase 3 trial, ORR was 80%, CR rate was 30%, and median time to progression was 11 months.⁸⁹ In relapsed/refractory MALT lymphoma, this approach resulted with ORR of 90–94%, CR rate of 62–77%, and median PFS exceeding 3 years.^{85,90} Therefore, ^{90}YIT may

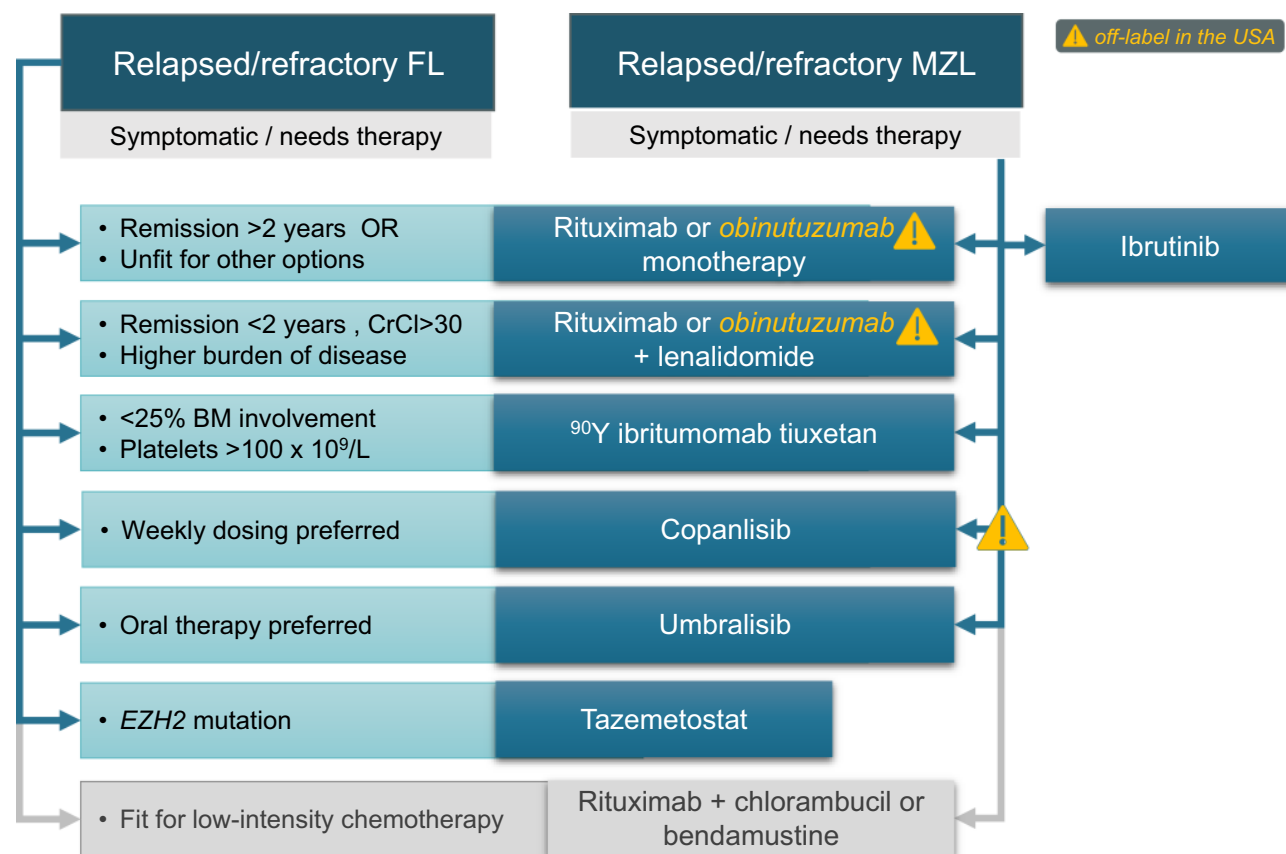


Figure 2 FDA-approved chemotherapy-free options for relapsed or refractory FL or MZL (note that some options are off-label in the USA for either one or both histologies).

Abbreviations: BM, bone marrow; CrCl, creatinine clearance; FL, follicular lymphoma; IV, intravenous; MZL, marginal zone lymphoma.

be an excellent option for an older patient with a relapsing FL/MZL and no prior exposure to chemotherapy.

Lenalidomide ± Rituximab or Obinutuzumab

Lenalidomide plus an anti-CD20 antibody offers an attractive approach for patients who can tolerate the associated toxicity, who did not receive lenalidomide as first-line therapy, and who have creatinine clearance > 30 mL/min. Similar to the first-line setting, this option is comparable in efficacy and toxicity to cytotoxic chemotherapy and may not be suitable for frail individuals who would otherwise not accept or withstand chemotherapy.

R2 is approved in both FL and MZL based on the phase 3 AUGMENT study which showed longer PFS (39 versus 14 months) and higher ORR (78% versus 53%) compared with rituximab.⁹¹ The median age of participants was 64 (range, 26–86), and 84% had prior rituximab exposure. Toxicity was unfortunately also higher with R2: 69% of patients experienced grade 3/4 toxicities (primarily

hematologic) and 26% had at least 1 serious adverse event. Neutropenia was the most common adverse event (58%), responsive to dose interruptions and reductions of lenalidomide which occurred in 64% and 26% of participants, respectively. It is important to note that the lenalidomide dose was 20 mg for patients with creatinine clearance >60 mL/min, but only 10 mg for those with clearance 30–59 mL/min. We suggest aggressive dose adjustments (including a potential start at 50% of the target dose with subsequent escalation) for older/unfit patients to avert the risk of toxicity-related hospitalizations and early discontinuations. In the phase 2 GALEN study for relapsed/refractory FL, lenalidomide with obinutuzumab also provided a high ORR (79%) and 2-year PFS (65%).⁹² However, adverse events were common with asthenia in 61%, neutropenia in 43%, diarrhea in 40%, bronchitis in 41%, and muscle spasms in 39%. While the advantage of obinutuzumab in this setting remains uncertain, responses may be higher for patients who had short (<2 years) remissions to prior rituximab-based therapy.

Table 2 Chemotherapy-Free Options for Relapsed/Refractory FL or MZL

Treatment	N	N FL	N MZL	Age, Median (Range)	Prior Lines, Median (Range)	Gr 3/4 Adverse Events	ORR	CR/CRu	Median PFS (mo)
Anti-CD20 monotherapy									
Obinutuzumab ⁴⁶	40	34		61 (42–79)	3 (1–11)	33%	55%	9%	12
Rituximab ⁴⁷		75	5	60 (38–80)	2 (1–6)	15%	33%	5%	25
Obinutuzumab ⁴⁷		74	6	62 (33–84)	2 (1–7)	15%	45%	12%	18
Rituximab ⁴³	219	214		62 (26–85)	2 (1–9)	28%	66%	20%	19
Ofatumumab ⁴³	219	214		61 (27–90)	2 (1–10)	37%	50%	16%	16
Rituximab ⁸⁹	70	58		57 (36–78)	2 (1–5)	^a	56%	20%	10 ^b
Ibritumomab tiuxetan									
⁹⁰ YIT ⁸⁹	73	55		60 (29–80)	2 (1–6)	≥ 60% ^a	80%	34%	11 ^b
⁹⁰ YIT ⁸⁵	30		30	57 (36–83)	2	≥ 60% ^a	90%	77%	NR
⁹⁰ YIT ⁹⁰	16		16	58 (19–77)	2	≥ 44% ^a	94%	63%	38
Lenalidomide ± anti-CD20 antibody									
Lenalidomide ⁹³	45	45		63 (34–85)	NR	58%	53%	20%	13 ^b
Lenalidomide ⁹⁴	43	22	3	63 (42–89)	3 (1–17)	NS	23%	5%	4
Lenalidomide + rituximab ⁹¹	178	147	31	64 (26–86)	1 (1–12)	69%	78%	34%	39
Lenalidomide + obinutuzumab ⁹²	88	88		64 (39–87)	2 (1–7)	≥ 44%	79%	39%	65% at 2 y
PI3K inhibitors									
Umbralisib ¹⁰³	208	117	69	66 (29–88)	2 (1–10)	53%	FL 45% MZL 49%	FL 5% MZL 16%	FL: 11 MZL: NR
Copanlisib ^{104,105}	142	104	23	63 (25–82)	3 (2–9)	≥ 53% ^a	FL 59% MZL 70%	FL 14% MZL 9%	11
Idelalisib ⁹⁸	125	72	15	64 (33–87)	4 (2–12)	54%	FL 54% MZL 47%	FL 6%	11
Duvelisib ¹⁰¹	129	83	18	65 (30–90)	3(1–18)	88%	FL 42% MZL 39%	FL 1% MZL 6%	9
BTK inhibitor									
Ibrutinib ¹⁰⁶	110	110		62 (28–87)	3 (2–13)	48%	21%	11%	5
Ibrutinib ¹⁰⁷	40	40		64 (46–82)	3 (1–11)	43%	38%	13%	14
Ibrutinib ¹⁰⁹	63		63	66 (30–92)	2 (1–9)	71%	58%	3%	16
EZH2 inhibitor									
Tazemetostat ¹¹¹ (EZH2 ^{mut})	45	45		62 (57–68)	2 (2–43)	27%	69%	13%	14
Tazemetostat ¹¹¹ (EZH2 ^{wt})	54	54		61 (53–67)	3 (2–5)	27%	35%	4%	11

Notes: ^aNot explicitly stated; ^btime to progression; ^cserious adverse events.

Abbreviations: AE, adverse event; ORR, overall response rate; CR/CRu, complete response/complete response unconfirmed; PFS, progression-free survival; mo, months; NR, not reached; y, years; ⁹⁰YIT, ibritumomab tiuxetan.

Lenalidomide (started at 15 mg during cycle 1, and escalated to 20 mg in cycles 2 to 12) has also been studied as a single agent, providing 53% ORR and 20% CR rate in recurrent FL with median time to progression of 1.1 year.⁹³ These results were inferior, and toxicity not lower, compared with R2, suggesting an important synergy with rituximab, which is consistent with the immunomodulatory mechanism of action for lenalidomide. ORR was even lower (23%) in a study enrolling various iBCL.⁹⁴ In the AUGMENT trial, no difference in the global quality of life assessment between R2 and lenalidomide was reported at any timepoint.⁹⁵ Therefore, we recommend using lenalidomide with an anti-CD20 antibody whenever possible. For elderly patients who find frequent visits to an infusion center burdensome and desire all-oral regimens, we favor other options discussed below.

New PI3K Inhibitors: More Specificity, Less Toxicity

The phosphatidylinositol 3-kinase (PI3K) inhibitors target the deregulated PI3K pathway important for B-cell receptor signaling, cellular proliferation, and anti-tumor immunity.⁹⁶ Currently, four PI3K inhibitors agents are approved by the FDA: one intravenous (copanlisib, approved for FL) and targeting predominantly PI3K α/δ , and three orally administered PI3K $\delta\pm\gamma$ inhibitors: idelalisib (FL, SLL/CLL), duvelisib (FL, SLL/CLL), and umbralisib (FL, MZL). Many clinicians find this class of drugs challenging because of differences between individual agents and because of severe infectious and autoimmune toxicity associated with some.⁹⁷ However, the novel PI3K δ inhibitor umbralisib appears to have improved toxicity, restoring an important lower-intensity option for older patients with FL or MZL.

Idelalisib was the first approved PI3K δ inhibitor based on a phase 2 trial that reported a 57% ORR in relapsed/refractory iBCL with median PFS of 11 months.⁹⁸ Toxicities included neutropenia (56%, grade ≥ 3 in 27%), diarrhea (43%, grade ≥ 3 in 13%), fatigue (30%, grade ≥ 3 in 2%), nausea (30%, grade ≥ 3 in 2%), cough (29%, no grade ≥ 3) and pyrexia (28%, grade ≥ 3 in 2%), as well as 13% grade 3/4 hepatitis. Serious toxicities were even more frequent in a real-world sample of older Medicare beneficiaries receiving idelalisib.⁹⁹ Longer use has amplified the safety concerns about colitis, hepatitis, pneumonitis, neutropenia, and opportunistic infections (CMV, *Pneumocystis jirovecii* pneumonia) and ultimately dissuaded many clinicians from routine use of

idelalisib.¹⁰⁰ Duvelisib was also studied in a phase 2 study of iBCL patients after median 3 lines of therapy. ORR was 47% and median PFS 9.5 months.¹⁰¹ Adverse effects were similar to idelalisib, including diarrhea (49%, grade ≥ 3 in 15%), neutropenia (29%, grade ≥ 3 in 25%), nausea (29%, grade ≥ 3 in 2%), fatigue (28%, grade ≥ 3 in 5%) and cough (27%, no grade ≥ 3). Grade 3 colitis and pneumonitis were seen in 8% and 5%, respectively, and a third of patients discontinued duvelisib because of toxicity.

Umbralisib is a novel drug in this class, characterized by substantially improved PI3K δ specificity and additional anti-casein kinase-1 ϵ (CK1 ϵ) activity which may attenuate the autoimmune toxicities.^{102,103} It is also the only oral PI3K inhibitor dosed once daily (800 mg), and has gained FDA approval for relapsed/refractory MZL and FL. In the registration phase 2 trial of 208 patients with iBCL (56% FL, 33% MZL) after median 2 prior therapies, the ORR was 45% in FL (5% CR) and 49% in MZL (16% CR), while median PFS was 11 months and not reached, respectively.¹⁰³ Rates of grade 3/4 diarrhea (10%), hepatitis (7%) and neutropenia (11%) appeared lower than with other PI3K inhibitors, and colitis and pneumonitis were very rare (any grade events: 1.9% and 1.4%, respectively). Considering this improved toxicity profile and convenience of once-daily dosing, umbralisib may be the PI3K inhibitor of choice for older or more frail patients. However, prophylaxis against *Pneumocystis jirovecii* and zoster, monitoring of CMV viremia, as well as strict dose reductions or holds are mandatory to maintain safety with this drug.

Copanlisib predominantly targets PI3K α and PI3K δ isoforms, resulting in a unique adverse event profile with transient hypertension, hyperglycemia, or neutropenia, but few long-term immune toxicities typical of PI3K δ/γ -targeting agents. It is approved for FL after ≥ 2 prior systemic therapies and has a breakthrough designation for MZL.^{104,105} In the phase 2 registration trial, the ORR in FL was 59%, CR rate was 14%, and median PFS was 11 months.¹⁰⁴ In MZL, copanlisib yielded ORR of 78% with median PFS of 24 months; CR (13%) occurred in SMZL.¹⁰⁵ Toxicity was substantial, including any grade fatigue in 52%, diarrhea in 48%, hyperglycemia in 48% (grade 3/4 in 39%), and hypertension in 44% (grade 3/4 in 39%), with 74% of patients requiring dose delays. Given the availability of oral alternatives, the use of copanlisib makes the most sense in the settings of financial barrier to accessing oral agents, or for patients who prefer weekly dosing.

Ibrutinib: For MZL Only

Although best known for their use in SLL/CLL and mantle cell lymphoma, Bruton tyrosine kinase (BTK) inhibitors have been evaluated in both FL and MZL.^{106–108} The efficacy in FL is notably low: in a larger phase 2 trial, the ORR was only 21% with median PFS of 5 months, though in a smaller study it reached 38%.^{106,107} In the latter experience, FL with *CARD11* mutations was resistant to ibrutinib, while duration of response was longer with *KMT2D* and *FOXO1* mutations.¹⁰⁷ If next-generation sequencing becomes routinely applied in relapsed/refractory FL, knowledge of *CARD11*, *KMT2D*, and *FOXO1* mutation status may allow a more personalized use of ibrutinib in FL. At present, single-agent BTK inhibitors in FL do not indicate sufficient efficacy.

In MZL, the use of ibrutinib is more compelling, as it attains 58% ORR (81% for those pre-treated with rituximab only) with a median PFS of 16 months and median duration of response of 28 months.^{108,109} It is now FDA-approved for use in relapsed/refractory MZL. Ibrutinib is tolerable among older patients, with the oldest patient enrolled in a phase 2 trial being 92 years old. Grade ≥ 3 AEs occurred in 71% of patients including grade ≥ 3 infection in 22%.¹⁰⁹ Most adverse events were grade 1/2, and grade ≥ 3 events included also anemia (16%), pneumonia (8%) and fatigue (6%). Ibrutinib is thus an excellent option for older patients whose MZL is refractory to rituximab monotherapy. Clinicians should be mindful of its potential cardiovascular toxicity (hypertension, atrial fibrillation), as well as interactions with anticoagulants and certain medications common in the elderly (diltiazem, verapamil). Ongoing studies explore the activity of other BTK inhibitors in MZL.

Tazemetostat: Personalized Therapy for FL with *EZH2* Mutation

Tazemetostat, an oral inhibitor of the enhancer of zeste homolog 2 (*EZH2*), has generated significant interest in FL, where the prevalence of activating *EZH2* mutations reaches 22%.¹¹⁰ Tazemetostat is approved in the US for FL with *EZH2* mutation, or for any FL without satisfactory options after ≥ 2 lines of systemic therapy. The reported ORR in the *EZH2*-mutated cohort from the phase 2 trial was 69%, including 13% rate of CR, whereas in *EZH2*-wild type these rates were 35% and 4%, respectively.¹¹¹ Median PFS was 14 and 11 months, respectively. As a targeted agent with high specificity, tazemetostat has an excellent toxicity

profile, with rare non-hematologic grade 3/4 adverse events (none more frequent than 3%). Grade 1/2 reactions are more common, including nausea (23%), diarrhea (18%), alopecia (17%), cough (16%), fatigue (15%) and upper respiratory infections (15%). Eligibility for tazemetostat can be determined using the FDA-approved single-gene *EZH2* mutation test or using a larger next-generation sequencing panel. For eligible older patients with FL, tazemetostat offers a particularly high-value therapeutic option.

Immunotherapy: Emerging Direction

Given the ongoing lack of curative options, many new agents are being studied for the treatment of iBCL (Table 3). Immunotherapy approaches are especially promising, as they offer novel mechanisms of action with high rates of CR and limited toxicity that is suitable for application among older patients. Both FL and MZL are known to strongly interact with the immune microenvironment, providing an opportunity to treat them by way of inducing immune attack or through immunomodulation within the tumor stroma.^{112,113}

Antibody-Drug Conjugates

Antibody-drug conjugates (ADC) combine a lymphoma-targeting monoclonal antibody with a cytotoxic small molecule to be internalized into the malignant cell upon binding. The goal is to increase tumor killing with minimal systemic toxicity. The first ADC established in lymphoma space, brentuximab vedotin, is unsuitable for FL or MZL which do not express its target CD30. Recently, ADCs targeting CD79b, CD22, or CD19 have been investigated in iBCL. Inotuzumab ozogamicin (which targets CD22) has been trialed in relapsed/refractory iBCL in a phase 2 study.¹¹⁴ The ORR was high (67%), but unfortunately so were the rates of adverse events, and 58% of patients discontinued therapy secondary to toxicity.

Polatuzumab vedotin is an ADC with a CD79b target conjugated with monomethyl auristatin E (MMAE), a microtubule inhibitor. It is FDA-approved in combination with BR for DLBCL and has been studied in other relapsed/refractory lymphomas.^{115,116} The phase 2 ROMULUS study compared the combination of rituximab with polatuzumab or with pinatuzumab vedotin (a CD22: MMAE conjugate) among patients with relapsed/refractory DLBCL and FL.¹¹⁷ Among 20 patients with FL receiving rituximab and polatuzumab vedotin, 70% achieved a response, including 45% with a CR, and

median PFS was 15 months. Grade ≥ 3 adverse events in FL were uncommon, including neutropenia (15%), diarrhea (10%) and dyspnea (5%). Polatuzumab vedotin is associated with cumulative peripheral neuropathy, ultimately leading to treatment discontinuation in 55% of patients with FL. Although more research on this agent is needed in iBCL to determine optimal dose and treatment duration, high efficacy and low toxicity make this drug potentially appropriate for older patients.

Loncastuximab tesirine is a novel anti-CD19 ADC investigated for aggressive and indolent lymphomas. Current experience is limited to a Phase 1 trial which enrolled 14 subjects with FL.¹¹⁸ The maximum tolerated dose was not reached, and ORR in FL was 79% with median duration of response not reached. The “naked” anti-CD19 antibody tafasitamab had a lower ORR in

a phase 2a trial (29% in FL), but it may synergize with lenalidomide as evidenced by its activity in DLBCL, which led to an FDA approval.^{119,120}

Chimeric Antigen Receptor (CAR) T Cell Therapy

Autologous CD19-directed CAR-T cells, while initially studied in B-lymphoblastic leukemia and DLBCL, appear to work quite well in iBCL may be even better tolerated than in the aggressive histologies.¹²¹ The ZUMA-5 phase 2 trial of axicabtagene ciloleucel included patients with FL and MZL who had received ≥ 3 lines of therapy, had performance status 0 or 1 and age up to 79.¹²² ORR was 92% (94% in FL and 85% in MZL) with a 76% CR rate (80% and 60%, respectively) and projected 12-month duration of response of 72%. The hallmark risks of CAR

Table 3 Selected Emerging Therapies in FL and MZL

Treatment	N	Age, Median (Range)	Prior Lines, Median (Range)	Gr. 3/4 Adverse Events	ORR	CR	Median PFS (mo)
Polatuzumab vedotin +R ¹¹⁷	20 (FL)	67 (59–74)	2 (2–4)	50%	70% ^a	45% ^a	15
Loncastuximab tesirine ¹¹⁸	14 (FL)	63 (20–87) ^b	3 (1–13)	77%	79% ^a	64% ^a	NR
Tafasitamab ¹¹⁹	34 (FL)	62 (40–87)	3 (1–>3)	27%	29%	9%	9
Axicabtagene ciloleucel ¹²²	124 (FL) 22 (MZL)	61 (34–79)	3 (1–10)	86%	FL 94% MZL 85%	FL 80% MZL 60%	NR
Blinatumomab ¹²⁴	38 (FL)	65 (20–80) ^b	3 (1–10)	90%	80% ^a	40% ^a	NR
Mosunetuzumab ¹²⁷	62 (FL)	59 (27–85)	3 (2–11)	35%	68%	50%	12
Epcoritamab ¹²⁹	12 (FL)	73 (35–84)	4.5 (1–18)	NR	100%	25%	NR
Nivolumab ¹³³	92 (FL)	67 (37–87)	3 (2–10)	15%	4%	1%	2
Atezolizumab + Obi orR ¹⁴¹	21 (MCL)	67 (47–87) ^b	2 (1–7)	48%	43%	14%	NR
Avadomide + Obi ^{140,142}	53 (FL)	61 (26–83) ^b	3 (2–4)	46%	76% ^a	47% ^a	17
Venetoclax ¹³⁷	29 (FL)	64 (46–75)	3 (1–10)	56%	38%	14%	11
Venetoclax +R ¹³⁸	52 (FL)	63 (40–84)	3 (1–6)	50%	35%	17%	7
Zanubrutinib ¹⁴³	68 (MZL)	70 (37–95)	2 (1–6)	29%	60%	15%	NR

Notes: ^aFL only; ^bfor all histologies.

Abbreviations: AE, adverse event; BR, bendamustine and rituximab; CR, complete response; mo, months; NR, not reported; Obi, obinutuzumab; ORR, overall response rate; PFS, progression-free survival; R, rituximab.

T-cell therapy, namely grade 3/4 cytokine release syndrome and neurotoxicity, were relatively reduced (7% and 19%, respectively) compared with observations from DLBCL. However, grade 3/4 adverse effects occurred overall in 86% of participants, including neutropenia in 33% and anemia in 23%. Based on these results, axicabtagene ciloleucel has been approved by the FDA for treatment of FL. However, due to the need for lymphodepleting chemotherapy and the overall intensity of the procedure, CAR T-cell therapy, while highly effective and potentially curative, will likely remain an option for only few older/unfit patients with iBCL. Experience with CAR T-cells among older (age 65–76) patients with DLBCL is so far encouraging, with similar efficacy and rates of cytokine release syndrome, although higher neurotoxicity than among younger patients.¹²³ However, the risk/benefit ratio of CAR T-cell therapy for iBCL in this population remains to be delineated.

Bispecific Antibodies

Bispecific antibodies are a novel class of immunotherapeutic agents which simultaneously bind an antigen (CD20 or CD19) on malignant B-cells, and CD3 on immune effector T-cells, thus facilitating the formation of the immune synapse and T-cell activation. They offer perhaps the most promising approach for relapsed/refractory and possibly also previously untreated iBCL. Their effects are similar to CAR T-cell therapy but without the need for lymphodepleting chemotherapy or apheresis. Their suitability for older and more frail patients who cannot tolerate cytotoxic chemotherapy has already been investigated in some early-phase trials. Blinatumomab (a CD19/CD3-directed bispecific T-cell engager) was studied in a phase 1 trial for patients (up to age 80) with relapsed/refractory FL and other lymphomas.¹²⁴ ORR among 15 patients with FL was 80% with 40% of patients attaining a CR, but the use of blinatumomab is complicated by the need for continuous intravenous infusion over weeks as well as 22% rate of grade ≥ 3 neurologic events.

Mosunetuzumab is a CD20/CD3 bispecific antibody which has been granted FDA breakthrough designation for FL.¹²⁵ The phase 1 trial of mosunetuzumab in relapsed/refractory CD20-positive lymphomas included 62 patients with FL of whom 68% responded and 50% achieved a CR.^{126,127} So far, 74% of patients who attained CR have remained in remission, and median duration of response is 20 months.¹²⁷ The average age of participants was 59, but ranged up to 85. One third of patients

experienced a treatment-related grade 3/4 adverse event, but they were mostly laboratory-based (neutropenia) and no grade 3/4 cytokine release syndrome or neurotoxicity occurred. Of note, mosunetuzumab has been studied specifically in the population of older and unfit patients with DLBCL (median age of 84, range 67–100), demonstrating a similar toxicity profile.¹²⁸ Due to its high efficacy and low toxicity, mosunetuzumab is moving forward with further testing in both relapsed/refractory and previously untreated FL/MZL. Other CD20/CD3 bispecific antibodies with ongoing studies in FL/MZL include epcoritamab,¹²⁹ glofitamab,¹³⁰ and odronextamab, but their safety in older patients will require more research.¹³¹

Checkpoint Inhibitors

Nivolumab as a single agent initially demonstrated promise with responses among 4 of 10 FL patients in a phase 1 study.¹³² Unfortunately, in the subsequent phase 2 experience, the ORR was only 4% with no clear predictive biomarker.¹³³ Single-agent PD-1 blockade may thus be ineffective in relapsed/refractory FL, although additional research aims to determine ways to enhance immune response or identify predictors of response.^{134,135} An intriguing case of a complete hematologic remission in SMZL after pembrolizumab has been reported, although otherwise data on checkpoint inhibitors in MZL are lacking.¹³⁶

Other Targeted Agents

Even though the *BCL2-IGH* rearrangement constitutes a molecular hallmark of FL, single-agent BCL2 inhibitor venetoclax is less active in FL than in SLL/CLL or mantle cell lymphoma. Among 29 patients in a phase 1 trial of venetoclax (with dose escalated up to 1200 mg) ORR was 38% and CR rate was 14%, with median PFS of 11 months.¹³⁷ The combination of venetoclax (800 mg daily) with rituximab has also been compared against the triplet of BR plus venetoclax or against BR in a phase 2 study for relapsed/refractory FL.¹³⁸ While the BR plus venetoclax demonstrated unacceptable toxicity, venetoclax with R had ORR of 35%, CR rate of 17%, and median PFS of 7 months, with a safety profile similar to that observed in CLL/SLL.^{138,139} These results were ultimately disappointing and indicate that further investigation of venetoclax in FL needs to identify patients who derive benefit from treatment with BCL2 inhibitors alone or in synergistic combinations.

Given the activity of lenalidomide, novel immunomodulatory imide drugs (IMiDs) are also under investigation. Avadomide combined with obinutuzumab shows similar tolerability to R2 with a high (76%) ORR in FL and 47% of patients achieving a CR.¹⁴⁰ Other IMiDs/cereblon E3 ligase modulators like iberdomide (CC-220) and CC-99282 are being further investigated to determine their activity in lymphoma.

Conclusions

First-line therapy for older patients with FL/MZL who are unfit for chemotherapy is still primarily based on rituximab monotherapy. Lenalidomide in combination with rituximab offers an alternative to chemotherapy with comparable efficacy, but toxicity is not sufficiently improved. In second and subsequent lines of treatment, novel therapies have changed the landscape, requiring a nuanced approach between FL and MZL subtypes, with many chemotherapy-free options possible for patients relapsing after anti-CD20 antibody therapy. The incoming wave of highly effective and relatively non-toxic immunotherapy approaches may further allow patients with iBCL to manage their disease without the need for chemotherapy exposure in their lifetime.

Disclosure

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