Radioimmunotherapy—A New Approach in the Management of Non-Hodgkin Lymphoma

a report by

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Despite the sensitivity of most lymphomas to initial therapy with chemotherapy or radiotherapy, the majority of patients with advanced non-Hodgkin’s lymphoma (NHL) eventually relapse and die of their disease.1 Furthermore, patients with advanced low-grade lymphomas remain incurable and their survival has not altered since the early 1960s.2 The introduction of antibody (mAb)-based therapy initially with Rituximab in the late 1990s and, more recently, with the conjugation of radioisotopes to mAb as part of radioimmunotherapy (RIT) has provided fresh hope for NHL patients that their prognosis can be improved. Rituximab has now become integrated in the treatment of most NHL and while the response rates of rituximab as a single agent remain modest, with complete response rates in single figures,3 combining rituximab with combination chemotherapy (CHOP) has shown a survival advantage when used in the treatment of aggressive lymphomas, and dramatic improvements in progression-free survival.4-7

To date, antibodies directed towards the CD20 antigen have dominated the field of mAb therapy and RIT of lymphoma. CD20 is highly expressed on mature B-cells and present on 95% of B-cell lymphomas.8 It is neither internalised nor shed from the cell surface and appears on binding mAb to initiate signalling and trigger apoptosis through a caspase-dependent pathway.9

What is Zevalin RIT?

Zevalin is composed of the monoclonal antibody ibritumomab, covalently bound to tiuxetan, a high-

affinity chelator for the radioisotope $^{90}$Y. Ibritumomab is a murine IgG1 kappa monoclonal antibody that specifically targets the CD20 antigen.\(^{10}\) The tiuxetan chelator creates a high-affinity, stable urea-type bond between the antibody and the radioisotope, to prevent the radioisotope dissociating and circulating around the body.\(^{11}\)

The Zevalin treatment consists of a pre-treatment of the rituximab. The regimen is delivered over seven to nine days on an out-patient basis. A typical course of treatment involves a one-day intravenous (IV) infusion of rituximab 250mg/m\(^2\); a second IV infusion of rituximab on day seven, eight or nine; immediately followed by an IV ‘slow push’ 10-minute infusion of $^{90}$Y labelled Zevalin. By labelling mAbs with beta-emitting radioisotopes, radiation can be targeted to the tumour increasing the potential for clinical response. Lymphomas are highly radiosensitive tumours and localized irradiation is the only treatment that potentially offers curative treatment for those patients with early-stage low-grade lymphomas.\(^{12}\) RIT has the potential advantage over other mAb directed therapy that it may kill tumour cells not directly targeted by the mAb via the ‘crossfire effect’. This can provide cell kill to adjacent, antigen negative tumour cells.\(^{13}\)

The unlabeled antibodies may bring about anti-tumor effects not only by recruiting the host immune system but also through inducing cell cycle arrest or apoptosis. Pre-clinical work indicates that intrinsic cytotoxicity of mAbs in RIT may be as important as its ability to effectively deliver targeted radiotherapy.\(^{14,15}\) This suggests that there may be a synergistic interaction between the mAb effector mechanisms and the radiation.

$^{90}$Yttrium (now licensed for use with Zevalin under the brand name Yttracis\(^{®}\)) offers a number of advantages over the most commonly used radioisotope in oncology practice, $^{131}$Iodine. $^{90}$Y is a pure beta emitter delivering higher energy radiation (2.3MeV v 0.6MeV) at a longer path length (5.3mm versus 0.8mm). This enhances the crossfire effect and may be advantageous in treating larger, less well-vascularized tumour nodules.\(^{11}\) The physical half-life is 64 hours, which matches the biological half-life of murine monoclonal antibody of Zevalin, and the absence of penetrating gamma emissions enables delivery as an out-patient.

Zevalin is dosed according to the patient’s body weight and baseline platelet counts. For patients with platelet counts $\geq$150,000/mm\(^3\), 5MBq/kg body weight is given, up to a maximum allowable dose of 1,200MBq. For patients with platelet counts of 100,000-149,000/mm\(^3\), Zevalin is dosed at 11MBq/kg, up to a maximum allowable dose of 1,200MBq.

**Clinical Experience with Zevalin RIT**

Zevalin RIT has been used for over 10 years and has emerged as a safe, effective and well tolerated therapy for relapsed ‘low-grade’ NHL.\(^{11}\) The majority of patients treated in the registration approval studies had relapsed follicular lymphoma. These published studies consistently show high rates of

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Overall and complete response rates in relapsed follicular lymphoma.

Even in previously heavily pre-treated patients who had received a median of four different chemotherapy regimens and who either did not respond to prior rituximab therapy or had disease progression within six months of therapy, the overall response rate of 74%, with 15% complete response was seen. A randomized controlled trial of Zevalin versus rituximab in relapsed or refractory low-grade or transformed follicular B-cell NHL was performed with the primary aim of demonstrating superior response rates of Zevalin over Rituximab. Seventy-three patients received two doses of rituximab 250mg/m² a week apart as pre-dosing followed by a single dose of Zevalin 0.4mCi/kg (15Mbq/kg). Seventy patients in the control arm received rituximab 375mg/m² weekly for four weeks. The overall response rate was 80% for the Zevalin group versus 56% for the rituximab alone group (p=0.002). Complete responses were 30% and 16% in the Zevalin and rituximab groups, respectively. Both regimens were well tolerated but, as expected, there was more myelosuppression in the RIT group.

These highly promising results demonstrated that Zevalin led to superior overall and complete response rates to those seen with Rituximab, and for patients with follicular lymphoma this translated into a significant improvement in the time to treatment failure. An interesting and potentially important finding to emerge from this study was the high level of activity of Zevalin in patients who had become refractory to chemotherapy, which was significantly greater than that seen with Rituximab. This data suggests the Zevalin offers hope for meaningful clinical responses even in patients who have become refractory to chemotherapy.

Perhaps the most impressive finding to emerge from these initial studies was that around 70% of the patients who achieve a CR remain in remission for years—some patients treated in the early studies are now in remission for more than five years after Zevalin treatment, with a median follow-up of almost four years (see Table 1).

An analysis of prognostic factors has confirmed that this remarkable durability of response is unlikely to be accounted for by patient selection as most of these durable remissions have been achieved in heavily chemotherapy pre-treated and chemo-refractory patients with validated poor prognostic factors, such as extensive prior therapy (one to nine regimens), bulky disease, high lactate dehydrogenase (LDH) and extranodal disease. Only disease bulk correlated with the overall response rate (<5cm) (89 patients objective response rate (ORR) 90% (p<0.001).

Examining the toxicity seen for patients treated in the Zevalin trials (n=261) indicates that 28% will experience grade 4 neutropenia and 8% will experience grade 4 thrombocytopenia. The non-hematological toxicity is extremely modest and even the hematological toxicity can be minimized if appropriate precautions are taken. Patients who have been heavily treated with chemotherapy, such as those

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SAFETY SUMMARY: In clinical trials (N=356), the most common adverse events were part of an infusion-related symptom complex. Reported Grade 3 and 4 cytopenias included: lymphopenia (40%), neutropenia (6%), leukopenia (4%), anemia (3%), and thrombocytopenia (2%). In addition, there have been a limited number of post-marketing reports of prolonged pancytopenia, marrow hypoplasia, and late onset neutropenia (defined as occurring 40 days after the last dose of Rituxan) in patients with hematologic malignancies.

In both clinical studies and post-marketing surveillance, there have been a limited number of reports of bronchiolitis obliterans and pneumonitis occurring post-Rituxan infusion. Additionally, hepatitis B virus (HBV) reactivation with fulminant hepatitis, hepatic failure, and death has been reported in some patients with hematologic malignancies treated with Rituxan.

WARNING: Fatal Infusion Reactions: Deaths within 24 hours of RITUXAN infusion have been reported. These fatal reactions followed an infusion reaction complex, which included hypoxia, pulmonary infiltrates, acute respiratory distress syndrome, myocardial infarction, ventricular fibrillation, or cardiogenic shock. Approximately 80% of fatal infusion reactions occurred in association with the first infusion. Tumor Lysis Syndrome (TLS): Acute renal failure requiring dialysis with instances of fatal outcome has been reported in the setting of TLS following treatment with RITUXAN. Severe Mucocutaneous Reactions: Severe mucocutaneous reactions, some with fatal outcome, have been reported in association with RITUXAN treatment. Please see brief summary of prescribing information on adjacent page.


Many patients who responded once to Rituxan responded again

Duration of first response 9.8 Months

Duration of response in retreatment 15.0 Months

Median duration of response: initial therapy vs retreatment (n=23) 1,3

Renewed hope

In clinical studies, of patients who had relapsed following a previous response to Rituxan therapy, 38% responded for about 15 months when retreated with Rituxan. This suggests that Rituxan retreatment responses may be even more durable than initial Rituxan responses. 1,3
**RITUXAN® (rituximab) BRIEF SUMMARY**

**Adverse Reactions:**

RITUXAN has been associated with hypertension, neoplasms (including lymphoma), neutropenia, and infections. In single agent clinical trials, patients treated with RITUXAN have experienced transfusion reactions, respiratory distress, hypotension, or hypertension during infusion. Care should be taken to ensure that the patient’s blood pressure is controlled and the infusion is slowly administered.

**Hypersensitivity Reactions:** RITUXAN has been associated with hypertension in patients with high levels of circulating malignant cells. In patients with high levels of circulating malignant cells, hypertension may cause or contribute to symptomatology. In many cases, hypertension is associated with other symptoms such as chest pain or dyspnea. In patients experiencing hypertension, additional administration of antihypertensive agents may be required.

**Cardiovascular:** Cardiac disorders have been reported in patients treated with RITUXAN. Cardiac disorders include hypotension, chest pain, cardiac tamponade, myocardial infarction, pericarditis, and atrial fibrillation. In patients with cardiac disease, caution is advised when administering RITUXAN. In patients with history of cardiac disease, additional monitoring may be required.

**Neoplasms:** Neoplasms have been observed in patients treated with RITUXAN. Neoplasms include lymphomas, angiosarcoma, and anaplastic thyroid carcinoma. Care should be taken to monitor for the development of new neoplasms in patients treated with RITUXAN. Additional monitoring may be required.

**Infections:** Infections have been reported in patients treated with RITUXAN. Infections include pneumonia, cellulitis, and sepsis. Infections may be severe and require hospitalization. Additional monitoring may be required.

**Fatality:** Fatality has been reported in patients treated with RITUXAN. Fatality may be associated with the development of complications such as infection, hypotension, or respiratory distress. In patients with compromised immune function, additional monitoring may be required.

**Miscellaneous:** Miscellaneous reactions include rash, skin infections, and injection site reactions. Additional monitoring may be required.

**Adverse Reactions:** Additional adverse reactions that may be associated with treatment include hypertension, hypotension, chest pain, atrial fibrillation, and sepsis. Additional monitoring may be required.

**Mucocutaneous Reactions:** Mucocutaneous reactions include Stevens-Johnson syndrome, toxic epidermal necrolysis, and drug reaction with eosinophilia and systemic symptoms. In patients with mucocutaneous reactions, additional monitoring may be required.

**Phlebitis:** Phlebitis has been reported in patients treated with RITUXAN. Phlebitis may be associated with the development of complications such as infection, hypotension, or respiratory distress. In patients with compromised immune function, additional monitoring may be required.

**Other:** Other reactions that may be associated with treatment include hypertension, hypotension, chest pain, atrial fibrillation, and sepsis. Additional monitoring may be required.

**Precautions:**

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with a reduced platelet count (<150 x 109/L), need to be given a reduced dose of radioactivity. With this in mind, Zevalin 0.3mCi/kg (11.1MBq/kg) proved to be both safe and efficacious.13–16

Although the results for single agent RIT are encouraging, the future is likely to involve integrating RIT into chemotherapy treatment protocols, and the current challenge for clinical investigators is to determine the optimal approach of integrating Zevalin RIT into chemotherapy schedules. Early data using Zevalin given to consolidate clinical responses following both shortened or full course chemotherapy look extremely promising.

Shipley et al. presented impressive data demonstrating the ability of Zevalin to convert partial response after brief chemotherapy to complete responses. In this study patients were treated with four weekly infusions of Rituximab followed by three cycles of R-CHOP followed eight weeks later by Zevalin. Of the 22 patients who had completed the protocol there was a 40% CR rate before Zevalin, which was converted to a 86% CR rate after Zevalin.17

The large European intergroup study in previously untreated follicular lymphoma, which randomized patients to Zevalin after initial chemotherapy, has now completed accrual with over 400 patients recruited, and will provide invaluable data as to whether Zevalin may have a role as consolidation after primary chemotherapy.

Clinical responses have also been observed for Zevalin in transformed follicular and relapsed de novo diffuse large B-cell lymphoma (DLBC). Small numbers of patients were treated as part of the initial phase I/II study and response rates of 58% with a 33% CR rate were found. A recent European phase II trial with over 100 patients older than 60 with relapsed DLBC has confirmed this impressive response rate in large numbers of patients with relapsed DLBC unsuitable for transplantation.18 Given this high single agent activity, clinical trials are now under way to integrate Zevalin into the front line treatment of DLBC and a large intergroup study with a randomisation for 90Y ibritumomab tiuxetan (Zevalin) after full course R-CHOP chemotherapy is under way.

The dose-limiting toxicity from RIT is myelosuppression with delayed thrombocytopenia and neutropenia occurring at around four to six weeks after therapy. The extent and duration of the myelosuppression appears to depend on bone marrow reserve (amount of previous chemotherapy, age of patient), degree of bone marrow infiltration, pharmacokinetics of the mAb, and the stability of the radioimmunoconjugate.

Higher myeloablative doses may, however, be delivered safely with support from an autologous stem cell transplant. Zevalin RIT is currently being extensively investigated as a component of high dose therapy and Zevalin have been added to high dose chemotherapy with the ‘standard’ conditioning regimen of BCNU, Etoposide, Ara-C and Melphalan (BEAM). The early results confirm the feasibility of this approach and these important studies are on-going.

**Conclusion**

In summary, Zevalin RIT offers an excellent treatment alternative in relapsed follicular lymphoma and provides an attractive alternative in second relapse after R-Chemo schedules for many patients. Promising results are emerging in the treatment of aggressive lymphoma and RIT is now being integrated into treatment schedules for DLBC including high dose therapy and ASCT. The future for RIT is therefore likely to involve integration into chemotherapy schedules, and the recent clinical results suggests that this type of approach offers great promise for the future treatment of lymphoma.