A phase 1/2 trial of ublituximab, a novel anti-CD20 monoclonal antibody, in patients with B-cell non-Hodgkin lymphoma or chronic lymphocytic leukaemia previously exposed to rituximab

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Summary

This phase 1/2 study evaluated the safety, pharmacokinetic behavior and anti-tumour activity of ublituximab, a unique type I, chimeric, glycoengineered anti-CD20 monoclonal antibody, in rituximab-relapsed or -refractory patients with B-cell non-Hodgkin lymphoma (B-NHL) or chronic lymphocytic leukaemia (CLL). Induction therapy (doses of 450–1200 mg) consisted of 4 weekly infusions in cycle 1 for NHL and 3 weekly infusions in cycles 1 and 2 for CLL. Patients received ublituximab maintenance monthly during cycles 3–5, then once every 3 months for up to 2 years. Enrolled patients with B-NHL (n = 27) and CLL (n = 8) had a median of 3 prior therapies. No dose-limiting toxicities or unexpected adverse events (AEs) occurred. The most common AEs were infusion-related reactions (40%; grade 3/4, 0%); fatigue (37%; grade 3/4, 3%); pyrexia (29%; grade 3/4, 0%); and diarrhoea (26%; grade 3/4, 0%). Common haematological AEs were neutropenia (14%; grade 3/4, 14%) and anaemia (11%; grade 3/4, 6%). The overall response rate for evaluable patients (n = 31) was 45% (13% complete responses, 32% partial responses). Median duration of response and progression-free survival were 9.1/2 months and 7.1/7 months, respectively. Ublituximab was well-tolerated and efficacious in a heterogeneous and highly rituximab-pre-treated patient population.

Keywords: anti-CD20 monoclonal antibody, B-cell lymphoma, LFB-R603, TG-1101, ublituximab.

Monoclonal antibodies (MAb) targeting CD20 have improved the outcome of patients with B-cell malignancies and represent a major advance in treatment (Dotan et al, 2010; Weiner et al, 2010; Singh et al, 2014). While the activity of single-agent rituximab has been modest, its addition to standard chemotherapy markedly improves overall response rates (ORR), progression-free survival (PFS) and overall survival (OS) across all subtypes of B-cell non-Hodgkin lymphoma (B-NHL) and chronic lymphocytic leukaemia (CLL) (Habermann et al, 2006; Van Oers et al, 2006; Hochster et al, 2009; Parikh & Wierda, 2010). Maintenance therapy with rituximab following immunochemotherapy induction significantly improved responses and PFS in previously untreated and relapsed follicular lymphoma (FL) (Van Oers...
et al, 2006; Nastoupil et al, 2014). However, complete responses (CRs) with rituximab are rare, and indolent lymphomas, such as CLL and FL, remain incurable diseases, highlighting the need for well-tolerated agents effective in the relapsed state (Khouri, 2006; Farren et al, 2015). The development of novel anti-CD20 MAbs with activity in rituximab-resistant disease represents a major advance in the care of these patients.

Rituximab depletes B cells through three primary mechanisms: complement-dependent cytotoxicity (CDC), programmed cell death and antibody-dependent cellular cytotoxicity (ADCC) (Oflazoglu & Audoly, 2010; Winiarska et al, 2011). Approximately half of all patients who are initially responsive to rituximab are refractory to retreatment (Davis et al, 2000). The mechanisms of rituximab resistance are hypothesised to revolve around loss of cell-surface CD20 by transcriptional down-regulation, selection of a CD20-negative clone, ‘shaving’ of the CD20 antigen/antibody complex, or internalisation of the antigen/antibody complex into the target cell (Beum et al, 2006; Revzani & Maloney, 2011; Winiarska et al, 2011; Vaughan et al, 2014). Efforts to overcome rituximab resistance have focused on the development of new CD20-targeted MAbs that: (i) are active in ‘low’ CD20-expressing tumours; (ii) bind to unique CD20 epitopes; and (iii) have increased ability to engage the innate immune system, particularly ADCC (Oflazoglu & Audoly, 2010; Winiarska et al, 2011).

Next-generation anti-CD20 MAbs have been engineered to have a variety of features that distinguish them from rituximab, many of which revolve around improving ADCC and CDC (Table S1). Ofatumumab, a second-generation, fully human anti-CD20 MAb optimised for high CDC activity, has shown efficacy as a single agent in CLL patients refractory to fludarabine and alemtuzumab (Wierda et al, 2010). Other approaches to enhance the interaction between anti-CD20s and their targets include changing the amino acid sequence (Bowles et al, 2006) or glycosylation pattern of the IgG in a way that enhances interaction with the Fc receptor (FcR) on effector cells (Dalle et al, 2011). Such modifications can enhance the in vitro efficacy and appear to be safe in clinical trials (Salles et al, 2012). Third-generation anti-CD20 MAbs are Fc domain-engineered antibodies designed to improve therapeutic activity by enhancing Fc-gamma receptor-IIIa (FcyRIIIa) interactions, improving ADCC activity (Winiarska et al, 2011). Obinutuzumab, a type II glycomodified MAb, has improved ADCC over rituximab, with less CDC than ofatumumab. It is a humanised IgG2 class MAB approved for the treatment of CLL (Cartron et al, 2014). Obinutuzumab is also unique in how it cross-links CD20, resulting in enhanced direct cell death.

Ublituximab is a type I, chimeric, glycoengineered anti-CD20 recombinant IgG1 MAB produced in the rat cell line YB2/0. Interestingly, ADCC activity is known to be dependent on the fucose content of oligosaccharides bound to anti-CD20-MAbs (Konno et al, 2012). Anti-CD20 MAbs with low fucose content exhibit higher ADCC activity, while antibodies with high fucose content have less ADCC (Konno et al, 2012). The low fucose content in ublituximab’s Fc region results in increased affinity for the FcγRIIIa (or CD16) (Le Garff–Tavernier et al, 2014, 2015). This increased affinity is associated with potent in vitro ADCC against B cells compared with both rituximab and ofatumumab, particularly against tumour cells that express low levels of CD20, such as CLL (De Romeuf et al, 2008; Bellon et al, 2011; Le Garff–Tavernier et al, 2011). Ublituximab differs from obinutuzumab in at least 3 distinct ways: (i) ublituximab is an IgG1 class MAb, which exhibits improved ADCC, in contrast to obinutuzumab, which is an IgG2 class MAB; (ii) ublituximab and obinutuzumab bind to distinctly different epitopes on the CD20 domain (Figure S1); and (iii) obinutuzumab is humanised, while ublituximab is chimeric. Murine xenograft models of FL and mantle cell lymphoma have demonstrated superior anti-tumour activity and marked tumour-growth delay with ublituximab compared with rituximab (Tourais Estaves et al, 2011). In a first-in-human phase 1 study of single-agent ublituximab in heavily treated CLL-patients, all of whom had relapsed following fludarabine-based therapy and 58% of whom were refractory to prior rituximab, ublituximab was well-tolerated and active at doses up to 450 mg per infusion (Cazin et al, 2011, 2013). Infusion-related reactions (IRRs), pyrexia, neutropenia and thrombocytopenia were the most prevalent adverse events (AEs), with an ORR of 45%.

This phase 1/2 trial (ClinicalTrials.gov NCT01647971) was performed to determine the maximum tolerated dose (MTD), safety and efficacy of single-agent ublituximab in patients with relapsed/refractory B-cell NHL or CLL previously treated with rituximab.

Methods

Study design

This was an open-label, phase 1/2 trial performed at 6 sites (Fig 1). Phase 1 employed a Fibonacci 3 + 3 dose-escalation design (flat doses of 450, 600, 900 and 1200 mg). Patients with CLL/small lymphocytic lymphoma (SLL) received treatment on days 1, 8 and 15 during cycles 1 and 2 (28-day cycles). In the absence of disease progression or unacceptable toxicity, patients received maintenance ublituximab on day 1 of cycles 3–5, then every 3 months for a maximum of 2 years. Patients with NHL received ublituximab weekly during cycle 1 (days 1, 8, 15 and 22), no dose in cycle 2, and then received maintenance infusions on day 1 of cycles 3–5, followed by a dose every 3 months for a maximum of 2 years (Fig 1). The CLL/SLL treatment schedule was selected based on pharmacokinetic/pharmacodynamic (PK/PD) results from the first-in-human phase 1 study indicating profound B-cell depletion on a weekly-times-8 schedule. Since this was the first human experience with ublituximab in NHL, the dosing schedule for patients with NHL was selected...
to mimic the schedule used by other anti-CD20 regimens, with the inclusion of maintenance infusions. Pre-infusion treatment with an oral antihistamine and steroids was required. Flat doses of ublituximab were escalated as follows: (i) 450 mg, (ii) 600 mg, (iii) 900 mg and (iv) 1200 mg. During the phase 2 study, the recommended phase 2 dose (RP2D) was administered on the same schedule as in the phase 1. All patients provided written informed consent. The study protocol was approved by an Institutional Review Board and was conducted in accordance with the principles of Good Clinical Practice and the ethical principles outlined in the Declaration of Helsinki.

Study objectives

The phase 1 objectives were to determine the dose-limiting toxicity (DLT), MTD, RP2D and safety profile. Secondary objectives included determination of ORR, PK profiles, PFS, duration of response (DOR) and time to response. A DLT was defined as any of the following occurring within cycle 1 and considered related to therapy: grade 4 thrombocytopenia, grade 4 febrile neutropenia, grade 3 or greater non-hematological AE, any grade 5 event or any persistent AE that delayed administration of the next dose by >14 days. The MTD was defined as the dose at which <2 patients experienced a DLT. CLL patients were enrolled in the 600 and 900 mg cohorts only and were evaluated for DLTs separately from NHL patients. The primary objective of phase 2 was determination of ORR. Secondary objectives included DOR, time to response, PFS and assessment of safety.

Inclusion criteria

Patients were required to have histologically confirmed relapsed or refractory B-NHL or CLL/SLL, prior rituximab treatment, an Eastern Cooperative Oncology Group (ECOG) performance status ≤2, and adequate organ/marrow function with baseline absolute neutrophil count >1 × 10^9/l and platelet count >50 × 10^9/l. ‘Rituximab-refractory’ was defined as progression on or within 6 months of rituximab treatment. ‘Relapsed’ patients were defined as those who progressed more than 6 months after rituximab treatment. Patients with primary central nervous system lymphoma were eligible.

Exclusion criteria

These included pregnancy; active hepatitis B or C, human immunodeficiency virus or other acute infections; uncontrolled concurrent illness; severe allergy to human/mouse antibodies; history of malignancy (except basal cell carcinoma, in situ carcinoma of breast or cervix treated surgically with curative intent, or any malignancy that had been in CR for least 5 years); and chemotherapy or radiation ≤3 weeks or stem cell transplant ≤3 months prior to study entry.

Patient evaluations

Blood samples for complete blood count and chemistry were collected prior to each infusion up to cycle 12 and post-infusion during cycles 1 and 5. Other evaluations included chest x-ray, computed tomography (at screening, end of cycle 2 and approximately every 12 weeks thereafter), and positron emission tomography (PET, optional). Efficacy analyses were performed for any patient with one post-baseline measurement. Safety analyses were based on all registered patients who received at least 1 dose of ublituximab.

Outcomes

Assessment of response was based on the International Working Group (IWG) criteria for NHL (Cheson et al., 2007) and CLL (Hallek et al., 2008). Response assessments occurred at week 8 and every 12 weeks thereafter. AEs were based on published criteria (Common Terminology Criteria for Adverse Events, Version 4.0, https://evs.nci.nih.gov/ftp1/CTCAE/CTCAE_4.03_2010-06-14_QuickReference_5x7.pdf). Duration of response, time to response and PFS were determined per standard IWG criteria (Cheson et al., 2007; Hallek et al., 2008).

Pharmacokinetic assessment

Serum concentrations of ublituximab were analysed at 3P Lab (University of Wisconsin, Carbone Comprehensive Cancer Center, Madison, WI, USA). Parameters evaluated included peak (Gmax) and trough concentrations, terminal half-life, area under the concentration curve (AUC) and systemic clearance.
Blood samples were collected pre-dose, 1 min prior to end of infusion (EOI) and 0.5, 6, 24, 72 and 168 h post-EOI on days 1 and 22 of cycle 1 (C1D1 and C1D22) and C5D1. Samples were also collected pre-dose, 1 min prior to EOI and 30 min and 6 h post-dose on days 8 and 15 of cycle 1. Pre-dose samples were also drawn prior to each maintenance infusion (cycles 3, 4, 6 and 9) for up to 1 year.

Ublituximab concentrations were evaluated by flow cytometry. The assay was validated over the range of 25–1500 ng/ml, with a correlation coefficient of >98% and per cent coefficient of variation (CV) ranging from 3.3% to 9.6%.

Standard PK parameters were determined using non-compartmental methods. C_{max}, and trough (C_{min}) concentrations were determined by inspection of each individual’s concentration-time curve. If two peaks occurred in the same collection period, the first peak was recorded as the C_{max}. Terminal disposition rate constants were estimated by linear regression analysis of the last two time points (72 and 168 h) of the log-concentration versus time. Terminal half-lives (t_{1/2}) were calculated by dividing 0.693 by the elimination rate constant. The AUC was calculated using the linear trapezoidal rule up to the last collection time point (AUC_{0-168 h}), then extrapolated to infinity. Systemic clearance was determined by dividing dose by AUC. Differences among the kinetic parameter variables were evaluated using an unpaired two-tailed t-test.

Statistical analysis

Up to 80 patients could be enrolled. If 8 or more patients responded, the null hypothesis of ORR ≤5% would be rejected and replaced by the hypothesis that the response rate for the treatment was ≥15%. The planned sample size provided 90% power at alpha <5%. Early termination of the study would occur if no responders were observed among the first 20 enrolled patients.

PFS was determined using the methods of Kaplan–Meier (Kaplan & Meier, 1958). All statistical analyses were performed using a two-sided hypothesis test at the 5% level of significance, using SAS (Cary, NC, USA) Version 9.2.

Results

Patients

Thirty-five patients were enrolled between August 2012 and June 2015, and 29 discontinued before completing the planned 2 years of treatment. Seventy-one per cent of patients had received ≥2 prior rituximab-based regimens, and 43% were considered rituximab-refractory (Table I).

Four patients discontinued prior to the first efficacy assessment and were not evaluable for response (2 for AEs not related to study drug; 1 for a serious AE [pneumonia]; and 1 patient withdrew consent). All 35 patients were evaluated for safety.

At the end of the study, 21/35 (60%) patients had discontinued treatment for progression, while 8/35 (23%) patients stopped treatment for other reasons [AE/serious AE (n = 3); withdrawal of consent (n = 1); or investigator/patient decision (n = 4)]. The remaining 6/35 (17%) completed all treatment.

Dose determination and treatment

No DLTs were observed, hence no MTD was identified (Table II). There was no significant difference in the overall number of AEs among the four dose cohorts. There appeared to be no difference in ORR between 900 and 1200 mg, with a slightly higher incidence of haematological AEs observed (grade 3 neutropenia, anaemia, and thrombocytopenia) at the 1200-mg dose level. Hence 900 mg was selected as the RP2D.

Infusion times averaged 4 h for the first administration, 3 h for the second and third, and decreased to an average of 90 min for the fourth and all subsequent doses. Day 1 infusions were split for CLL patients with up to 150 mg administered on day 1 and the remaining dose administered on day 2 of cycle 1 only. Dose interruptions were permitted at the discretion of the investigator.

Safety

All 35 patients experienced ≥1 AE, with 17 (49%) reporting at least 1 grade 3/4 AE. Seventeen patients (49%) required ≥1 dose interruption, of which 11 were for IRRs. IRRs occurred in 14 (40%) patients (Table III) and were more prevalent among patients with CLL. The majority of IRRs occurred on C1D1, with only 5 occurring on subsequent cycles. No episodes of grade ≥3 IRR were reported. All IRRs were manageable with infusion interruptions, and all patients recovered without repercussion.

Other AEs included fatigue, pyrexia and diarrhoea (Table III). Laboratory abnormalities included neutropenia (14%; grade 3/4, 14%), thrombocytopenia (6%; grade 3/4, 6%) and anaemia (11%; grade 3/4, 6%). No infections were associated with grade 3/4 neutropenia, and no bleeding accompanied thrombocytopenia. All patients with grade 3/4 neutropenia received granulocyte colony-stimulating factor (G-CSF) and recovered without sequelae. No events of febrile neutropenia were reported. Urinary tract infection was reported in 5 patients, with only 1 grade 3/4 event. One event of hypogammaglobulinaemia was reported at the lowest dose studied (450 mg).

Serious AEs occurred in 13 patients (37%), the most frequent being pneumonia (n = 3). All other serious AEs occurred only once. No deaths were related to study treatment. One patient developed grade 3 serum sickness after 2 infusions, was hospitalised, permanently discontinued ublituximab, and recovered 15 days after the event was reported.
Efficacy

The best ORR to ublituximab among 31 patients evaluable for efficacy was 45% (Fig 2A). Forty-five per cent of patients had stable disease (SD), and 10% experienced progressive disease (PD) at the first efficacy assessment. Five patients experienced improvement in their response during the maintenance phase as follows: three patients (one patient each with CLL, marginal zone lymphoma [MZL] and FL) improved from SD to partial response (PR); while two patients (one patient each with FL and MZL) improved their response from PR to complete response (CR). Only 15 (48%) patients progressed during the maintenance phase, supporting the potential merits of protracted dosing in this heavily treated population.

The ORR was 44% (11/25) and 50% (3/6) in patients with NHL and CLL, respectively (Fig 2A). Among the 6 evaluable CLL patients, 5 (83%) had an absolute lymphocyte count (ALC) >4 × 10^9/l at study entry (range 3.055–165.996 × 10^9/l). A rapid depletion in circulating lymphocytes was observed in all CLL patients, with most patients achieving a >50% reduction within 7 days of the first infusion, and all 6 patients achieved an ALC <4 × 10^9/l within the first cycle (Fig S2). Of the 13 evaluable patients who were rituximab-refractory, 4 achieved an ORR of 31%, including 2 CRs, all in patients with indolent NHL. Change in tumour size from baseline is shown in Fig 2B.

The median time to response was 1.8 (range 1–11) months, with a median DOR of 9.2 (range 1–24) months. Median PFS for all evaluable patients was 7.7 (95% CI: 4.5–16.2) months with a median PFS for the rituximab-refractory patients of 4.7 (95% CI: 1.9–16.2) months (Fig 2C).

Pharmacokinetic profile

Twenty patients had PK samples submitted and analysed (Table SII). Sixteen patients were evaluable for full PK analyses during C1D1, 14 during C1D22, and 10 during C5D1. Figs 3A and 3B present the free drug concentration as a function of time curves, showing a dose-proportional relationship. There was a good relationship between dose and

| Characteristic | Phase I (N = 20) | Phase II (N = 15) | Patients (N = 35) |
|---------------|-----------------|------------------|-------------------|
| Median age – years (range) | 65 (50–88) | 69 (45–86) | 66 (45–88) |
| Gender – n (%) |                |                  |                   |
| Female       | 8 (40)          | 10 (67)          | 18 (51)           |
| Male         | 12 (60)         | 5 (33)           | 17 (49)           |
| ECOG – n (%) |                |                  |                   |
| 0            | 9 (45)          | 4 (27)           | 13 (37)           |
| 1            | 10 (50)         | 10 (67)          | 20 (57)           |
| 2            | 1 (5)           | 1 (6)            | 2 (6)             |
| Subtype of lymphoma – n (%) |            |                  |                   |
| Indolent NHL | 10 (50)         | 10 (67)          | 20 (57)           |
| Follicular   | 7 (35)          | 5 (33)           | 12 (29)           |
| Marginal zone | 3 (15)         | 5 (33)           | 8 (23)            |
| CLL/SLL      | 8 (40)          | –                | 8 (23)            |
| Aggressive NHL | 2 (10)        | 5 (33)           | 7 (20)            |
| Mantle Cell  | 2 (10)          | 3 (20)           | 5 (14)            |
| Diffuse Large B-Cell | –           | 2 (13)           | 2 (6)             |
| Prior therapy regimens – median (range) | 3.5 (1–6) | 2 (1–9) | 3 (1–9) |
| Prior therapy – n (%) |            |                  |                   |
| Rituximab    | –               | –                | 35 (100)          |
| Alkylating Agent (R-CHOP, R-CVP, R-ICE, other) | – | – | 23 (66) |
| Bendamustine (± rituximab) | – | – | 12 (34) |
| Purine analogue | –             | –                | 10 (29)           |
| Stem-cell transplantation | – | – | 5 (14) |
| Bortezomib   | –               | –                | 5 (14)            |
| Experimental therapy* | – | – | 6 (17) |
| Rituximab-refractory – n (%) | 7 (35) | 8 (53) | 15 (43) |
| 2 or > prior rituximab regimens – n (%) | 14 (70) | 11 (73) | 25 (71) |
| Refractory to immediate prior therapy – n (%) | 7 (35) | 8 (53) | 15 (43) |

CLL, chronic lymphocytic leukaemia; ECOG, Eastern Cooperative Oncology Group; NHL, non-Hodgkin lymphoma; R-CHOP, rituximab, cyclophosphamide, hydroxydaunorubicin (doxorubicin), Oncovin (vincristine) and prednisone; R-CVP, rituximab, cyclophosphamide, vincristine and prednisone; R-ICE, rituximab, ifosfamide, carboplatin and etoposide; SLL, small lymphocytic lymphoma.

*Includes bevacizumab, vorinostat, MLN4924, brentuximab, pralatrexate, lenalidomide.
喉部刺激。

†包括潮红、瘙痒、呼吸困难和全身发冷。

事件被分类到相同的预设术语类别中，计算一次。

在MedDRA版本18（http://www.meddra.org/how-to-use/support-documentation）中根据预设术语进行分类。

表II. Ublituximab活动与剂量群体（1期）

| Dose (mg) | Disease | DLT | Best response |
|----------|---------|-----|---------------|
| 450      | MZL     | No  | CR            |
|          | FL      | No  | PD            |
|          | MCL     | No  | PD            |
| 600      | FL      | No  | SD            |
|          | FL      | No  | SD            |
|          | MZL     | No  | CR            |
|          | CLL     | No  | NE            |
|          | CLL     | No  | NE            |
|          | CLL     | No  | PR            |
|          | CLL     | No  | PR            |
|          | FL      | No  | CR            |
|          | FL      | No  | CR            |
|          | MCL     | No  | SD            |
| 900      | FL      | No  | SD            |
|          | MZL     | No  | CR            |
|          | CLL     | No  | SD            |
|          | CLL     | No  | SD            |
| 1200     | FL      | No  | SD            |
|          | FL      | No  | PD            |
|          | MCL     | No  | SD            |

CLL, 慢性淋巴细胞白血病；CR, 完全响应；FL, 滤泡性淋巴瘤；MCL, 骨髓性淋巴瘤；MZL, 边缘区淋巴瘤；NE, 无法评估；PD, 进展性疾病；PR, 部分响应。

表III. 治疗期间的不良事件

| 不良事件（N = 35） | 任何级别 n (%) | 3/4级 n (%) |
|-------------------|----------------|------------|
| 冲击相关反应†     | 14 (40)        | 0          |
| 疲劳              | 13 (37)        | 1 (3)      |
| 头晕              | 10 (29)        | 0          |
| 腹泻              | 9 (26)         | 0          |
| 呕吐              | 8 (23)         | 0          |
| 失眠              | 6 (17)         | 0          |
| 呕吐               | 6 (17)         | 0          |
| 腹泻              | 5 (14)         | 1 (3)      |
| 脱水              | 5 (14)         | 5 (14)    |
| 周围水肿          | 5 (14)         | 1 (3)      |
| 慢性尿道感染      | 5 (14)         | 1 (3)      |
| 腹痛              | 4 (11)         | 0          |
| 减少的血红素      | 4 (11)         | 2 (6)      |
| 寒战              | 4 (11)         | 0          |
| 鼻窦炎            | 4 (11)         | 0          |

*不良事件发生在≥10%的患者中。

†冲击相关的反应包括寒战、瘙痒、呼吸困难和喉部刺激。

\( C_{\text{max}} \) (R2 = 0.69, Fig S3A) 和剂量和暴露的（AUC0-C1, R2 = 0.38, Fig S3B）对于C1D1中的ublituximab的剂量范围进行了研究。

讨论

将anti-CD20抗体引入B细胞恶性肿瘤的治疗中，为患有NHL和CLL的患者带来了改善的临床结果。但是，对rituximab的获得性耐药是一个重要的临床问题。刚成为或耐药的B细胞恶性肿瘤患者需要新的生物制剂来克服事先获得的rituximab耐药。

在一期试验中建立了ublituximab的安全性。最常见不良事件是1/2级IRR，没有3/4级IRR。在一期和二期试验中，obinutuzumab在15%和25%的CLL患者中表现出3/4级IRR。相比之下，ublituximab在90%的患者中出现IRR，但下降到平均90 min的多次给药后出现IRR。按照临床前研究，与other anti-CD20 MAb（https://www.pharma.us.novartis.com/sites/www.pharma.us.novartis.com/files/arzerra.pdf），乌力古单抗的毒性特征有利于或至少与用于其他anti-CD20 MAb相似。

Ublituximab在既往治疗的患者中产生了有意义的响应。对于31例既往治疗的患者和31%的13例rituximab-难治性患者。因为面临的挑战是在获得标准成像之前与研究进行，我们定义了rituximab-难治性人群作为难治性或在6个月内复发的患者。ublituximab和rituximab难治性组的响应率分别优于5%和44%的患者。UBIT与另一anti-CD20 MAb ofatumumab（https://www.pharma.us.novartis.com/sites/www.pharma.us.novartis.com/files/arzerra.pdf）相比，具有相似或至少相似的毒性特征。

虽然在特定亚组的比较上进行困难，但 ublituximab在CLL患者中的ORR为45%，尽管在13例ribitumab难治性患者中的ORR为31%。在13例ribitumab难治性患者中，ublituximab的ORR为45%，在31例可评估患者中为30%，在13例ribitumab难治性患者中为42%。与other anti-CD20 MAb比较，ublituximab的ORR为55%，44%的ORR出现在另一anti-CD20 MAb ofatumumab中。UBIT与另一anti-CD20 MAb ofatumumab（https://www.pharma.us.novartis.com/sites/www.pharma.us.novartis.com/files/arzerra.pdf），ublituximab的ORR为45%。
Fig 2. Treatment Response. (A) Overall response by lymphoma subtype. (B) Individual patient best percentage change from baseline in nodal size. (C) Progression-free survival. aNHL, aggressive non-Hodgkin lymphoma; CL, confidence interval; CLL, chronic lymphocytic leukaemia; CR, complete response; DLBCL, diffuse large B-cell lymphoma; FL, follicular lymphoma; iNHL, indolent non-Hodgkin lymphoma; MCL, mantle cell lymphoma; MZL, marginal zone lymphoma; ORR, overall response rate; PD, progressive disease; PFS, progression-free survival; PR, partial response; SD, stable disease.
provide meaningful clinical benefit to patients with limited treatment options.

Even considering the variability in sample size across subtypes, notable activity was observed in patients with indolent lymphoma, with 21% (4/19) of patients achieving a CR as their best response. In comparison, in the phase 2 obinutuzumab trials, CRs were achieved by 11% to 29% of patients with indolent lymphoma (Salles et al., 2013; Sehn et al., 2015). CR rates were lower with rituximab monotherapy (3–17%) (Feuring-Buske et al., 2000; Foran et al., 2000; Cohen et al., 2003; Sehn et al., 2015). Similarly, rituximab monotherapy has shown modest activity in CLL (Huhn et al., 2001; Furman et al., 2014), which has been attributed to low CD20 antigen expression by CLL cells (Prevodnik et al., 2011). The collective experience with ublituximab supports improved ADCC and improved activity in low CD20-expressing malignancies (De Romeuf et al., 2008; Bellon et al., 2011).

Anti-CD20 therapy has demonstrated the greatest benefit in combination, traditionally with multi-drug chemotherapy-based regimens. While the introduction of novel targeted therapies has shifted the treatment paradigm of CLL and indolent lymphoma, the activity of these agents is likely to be potentiated by the addition of an anti-CD20 MAb given their different mechanisms of action. Recently, a number of multi-drug, non-chemotherapy-based regimens have emerged, which although they are in early stages, appear to offer the efficacy of standard chemotherapy without the toxicity (Furman et al., 2014; Burger et al., 2015; Jones et al., 2015).

In similar fashion, ublituximab is being evaluated for the treatment of NHL or CLL in combination with other agents, including lenalidomide (ClinicalTrials.gov NCT01744912), ibrutinib (ClinicalTrials.gov NCT02013128), and TGR-1202 (PI3K-inhibitor) with or without ibrutinib (ClinicalTrials.gov NCT02006485). A phase 3 study will compare treatment with ibrutinib versus ibrutinib plus ublituximab in patients with previously treated CLL who have high-risk cytogenetic features (ClinicalTrials.gov NCT02301156).

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Author contributions

DM, JGK, PS, and HPM were responsible for the conception and design of the study. Collection and assembly of data was done by AS, CMF, MTS, MYK, DM, CD, JEA, PGN, JMK, PS, HPM, and OAO. Data analysis and interpretation were performed by CMF, DM, JGK, PS, HPM, and OAO. Manuscript writing was accomplished by all authors, and all authors reviewed and approved the final manuscript.

Conflicts of interest

MTS, CD, and OAO have received research funding from TG Therapeutics, Inc. (TGTX). OAO has also received funding for travel, accommodations, and expenses from TGTX, for which he has served as an unpaid consultant. PS and HPM are employees of TGTX who hold leadership roles and stock in the company. PGN has served as a consultant and received travel and accommodations expenses from Novartis and Boehringer Ingelheim (BI); he has received research funding from Novartis, BI, Genentech, TGTX, Lilly Newlink, Agendia, Abraxis, AstraZeneca, and Celgene. AS has received honoraria from Gilead Sciences. DM has been a speaker and received reimbursement for travel, accommodations, and expenses from Janssen and Alexion. JEA has received research funding from Acetylon. JMK has stock or other ownership to disclose with Helix Diagnostics. JGK has been a speaker for and received honoraria from Amgen, and has held a paid consulting or advisory role with TGTX. CMF and MYK have no conflicts of interest to declare.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Fig S1. Epitope binding sites on anti CD20 MAbs.
Fig S2. Change from baseline in absolute lymphocyte count (ALC) in patients with chronic lymphocytic leukemia (CLL).
Fig S3. Pharmacokinetic analysis. A. C_{max} vs. dose. B. Exposure versus dose.
Table S1. Differences between selected anti-CD20 monoclonal antibodies.
Table SII. Ublituximab pharmacokinetic summary.

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