Btk‐specific inhibition blocks pathogenic plasma cell signatures and myeloid cell–associated damage in IFNα‐driven lupus nephritis

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Introduction

Systemic lupus erythematosus (SLE) is often associated with exaggerated B cell activation promoting plasma cell generation, immune‐complex deposition in the kidney, renal infiltration of myeloid cells, and glomerular nephritis. Type‐I IFNs amplify these autoimmune processes and promote severe disease. Bruton’s tyrosine kinase (Btk) inhibitors are considered novel therapies for SLE. We describe the characterization of a highly selective reversible Btk inhibitor, G‐744. G‐744 is efficacious, and superior to blocking BAFF and Syk, in ameliorating severe lupus nephritis in both spontaneous and IFNα‐accelerated lupus in NZB/W_F1 mice in therapeutic regimens. Selective Btk inhibition ablated plasmablast generation, reduced autoantibodies, and — similar to cyclophosphamide — improved renal pathology in IFNα‐accelerated lupus. Employing global transcriptional profiling of spleen and kidney coupled with cross‐species human modular repertoire analyses, we identify similarities in the inflammatory process between mice and humans, and we demonstrate that G‐744 reduced gene expression signatures essential for splenic B cell terminal differentiation, particularly the secretory pathway, as well as renal transcriptional profiles coupled with myeloid cell–mediated pathology and glomerular plus tubulointerstitial disease in human glomerulonephritis patients. These findings reveal the mechanism through which a selective Btk inhibitor blocks murine autoimmune kidney disease, highlighting pathway activity that may translate to human SLE.
with glomerular cytokine/chemokine production, complement activation, and extensive proteinuria (4, 5). In NZB/W_F1 SLE–prone mice, direct activation of Fcγ receptor–bearing (FcγR-bearing) myeloid cells, including monocytes/macrophages, by glomerular ICs is sufficient to initiate inflammatory responses, resulting in tissue damage (6). The autoantibody IC also activate TLRs 7 and 9 in myeloid cells and plasma-cytoid DCs, leading to the secretion of IFNα that amplifies immune responses and consequently worsens disease (7, 8). IFNα augments B cell abnormalities in conjunction with TLR stimulation by lowering the activation threshold of autoreactive B cells, enhancing their survival and differentiation into plasmablasts and thereby triggering an excessive germinal center (GC) response (1, 2, 9, 10). In human SLE patients, enhanced IFNα stimulation, demonstrated through an IFN gene signature in blood, correlates with disease severity and higher ANA levels (11). Studies in NZB/W_F1 mice have confirmed the enhancing function of type-I IFNs in lupus pathogenesis. NZB/W_F1 mice deficient in type-I IFN receptor show prolonged survival (12), and conversely, adenovirus-mediated delivery of IFNα accelerates lupus manifestations, leading to severe glomerulonephritis (5, 13, 14).

Current treatments for severe SLE or LN, such as mycophenolate mofetil or cyclophosphamide (CTX), are effective at reducing mortality but fail to cure, and they are accompanied by severe adverse effects via their immunosuppressive or cytotoxic properties, respectively (15, 16). The only targeted immunotherapy approved for SLE is the anti-BAFF Ab belimumab that acts by reducing naïve and transitional B cells (17). However, initial clinical trials were not designed to assess the efficacy of belimumab for the treatment of LN. B cell depletion through anti-CD20 treatment has been studied in lupus, substantiating pathogenic roles of B cells, but clinical trials of anti-CD20 in SLE and LN have not supported approval (2, 9). Therefore, there is a high unmet need for targeted therapy in SLE. Because of the complexity of B cell involvement in disease pathogenesis, a drug that antagonizes more than one effector pathway would hold great therapeutic potential for more severe disease.

Bruton’s tyrosine kinase (Btk) is a Tec-family kinase that is expressed in most hematopoietic cells but not T cells. Btk is a key mediator of B cell receptor (BCR) signaling in B cells and FcγR signaling in myeloid cells (18–20). Mutations in the Btk gene lead to B cell deficiency manifested as X-linked agammaglobulinemia in humans and the related but less severe X-linked immunodeficiency in mice, emphasizing its role in B cell development. In animal models of arthritis, Btk inhibition abrogates both B cell– and myeloid cell–mediated disease marked by reductions in autoantibody and inflammatory cytokine levels (18, 21, 22). Furthermore, Btk deficiency and Btk inhibitors such as RN486, M7583, BI-BTK-1, and PF-06250112 of divergent selectivity profiles have shown benefit in preclinical models of SLE (23–30).

Given the fundamental role of Btk in both B cell and myeloid cell function and the chronic nature of treatment in autoimmune disorders, highly selective Btk inhibition is an attractive therapeutic strategy for SLE. Here, we show that specific Btk inhibition with a new prototype noncovalent, reversible Btk inhibitor G-744 — which exhibits over 400-fold selectivity against 284 kinases tested — abrogates aggressive disease in IFNα-accelerated NZB/W_F1 SLE and is superior to BAFF blockade or Spleen tyrosine kinase (Syk) inhibition. Furthermore, transcriptional profiling and modular expression analyses of spleen and kidney revealed that Btk inhibition, but not BR3-Fc or a Syk inhibition, abrogates gene signatures associated with plasma cell (PC) generation in the spleen, myeloid cell activation in the kidney, and decreased the expression of renal ortholog genes previously identified to show enhanced expression in the glomerular and tubulointerstitial compartments in human LN patients.

Results

G-744 is a noncovalent, highly selective small molecule inhibitor for Btk. We previously reported CGI-1746 as a highly selective, noncovalent, reversible inhibitor of Btk (18). Further lead optimization efforts identified the noncovalent, reversible Btk inhibitor G-744 (Figure 1A) (31) that had a similar binding mode and selectivity profile as CGI-1746 but improved physiochemical, absorption, distribution, metabolism, and excretion properties, allowing for oral dosing. In vitro enzyme assays, G-744 potently inhibited Btk kinase activity with an 50% inhibitory concentration (IC50) of 1.28 nM at apparent Km ATP concentration (Table 1). G-744 demonstrated > 1,000-fold biochemical selectivity over all 284 kinases tested except for EphA7 and Fgr, which still showed robust selectivity of 428-fold and 868-fold, respectively (Table 1 and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.90111DS1). We compared G-744 with other Btk inhibitors and showed that G-744 exhibited the highest selectivity against Tec-family kinases (Itk, Tec) and Src-family kinases (Blk, Frk, Hck, Lck, Lyn,
Src, and Yes), as well as several other kinases including Csk, EGFR, Flt3, and Jak3 (Supplemental Table 1). These results demonstrated that G-744 is a Btk inhibitor with a favorable selectivity profile that allows for a direct examination of the contribution of Btk to cellular functions and autoimmune disease pathogenesis independent of coinhibition of other enzymes, including Tec- and Src-family kinases.

**G-744 inhibits murine B cell and macrophage signaling.** We evaluated the cellular potency and functional activity of G-744 and compared it with a selective Syk inhibitor P505-15 (32) (Supplemental Table 1), since both Btk and Syk play a critical role in signal transduction of BCR and FcR (21, 33). G-744 potently inhibited anti-IgM–induced Btk-Y223 tyrosine phosphorylation (pBtk) in murine B cells (Figure 1B and Table 2) with an IC$_{50}$ of 20 nM. As expected, Btk inhibition had no effect on BCR-induced Akt phosphorylation on S473 (Figure 1C) (34). G-744 abrogated anti-IgM–induced Ca$^{2+}$ flux (Figure 1D) and murine B cell proliferation (Figure 1F and Table 2). Upregulation of the T cell costimulatory molecule CD86 on B cells after

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**Figure 1. G-744 inhibits Btk signaling and B cell receptor–mediated (BCR-mediated) responses in murine B cells.** (A) Chemical structure of G-744. (B–F) B cell signaling readouts in presence of indicated concentrations of G-744 or P505-15 following anti-IgM F(ab)$^{\prime}$2 stimulation of primary murine splenic B cells. (B) Phosphorylation of Btk-Y223 in presence of G-744. The triangle indicates pBtk-Y223 phosphorylation in unstimulated B cells without G-744. (C) Phosphorylation of Akt-S473 in presence of G-744. (D and E) Induction of [Ca$^{2+}$] in presence of (D) G-744 or (E) P505-15. Arrows indicate stimuli added. (F) Proliferation in presence of G-744. The triangle and square indicate B cell proliferation without G-744 in un- and anti-IgM F(ab)$^{\prime}$2-stimulated B cells, respectively. (G) Inhibition of anti-IgD–induced CD86 induction in C57BL/6 mice by G-744. CD86 expression on splenic B220$^+$ B cells was quantified by flow cytometry 6 h after anti-IgD stimulation. Mice were dosed orally with vehicle or G-744 as indicated prior to anti-IgD injection. Mean fluorescence intensity (MFI) is indicated for individual mice (represented by symbols), and data is expressed as mean ± SEM for each group (n = 5). Percent inhibition by G-744 treatment compared with stimulated vehicle control is indicated. *P < 0.05, **P < 0.0001, one-way ANOVA with Dunnett’s correction. (B–G) Graphs are representative of at least 3 experiments.
BCR activation in vitro was inhibited with similar potency (Table 2) and was significantly decreased in vivo compared with vehicle at concentrations of 10 mg/kg and above (Figure 1G). Furthermore, Btk inhibition abolished anti-CD40–stimulated B cell proliferation (Table 2). Syk inhibition with P505-15 blocked S473 Akt phosphorylation with similar potency as preventing B cell proliferation, consistent with a nonredundant

Figure 2. Specific Btk inhibition is superior to Syk or BAFF blockade in improving disease in IFNα-enhanced NZB/W_F1 LN. Fifteen-week-old NZB/W_F1 female mice (n = 10/group) were administered 1 injection of adenovirus containing murine IFNα (AdIFNα). Drug treatment started 3 weeks after AdIFNα injection with 100 mg/kg G-744 (triangles, dark-blue line), 60 mg/kg P505-15 (squares, red line), vehicle Hydroxypropylmethyl cellulose (HPMC) (open triangles, black line), 7.5 mg/kg BR3-Fc (circles, green line), Ig control antibodies (open circles, grey line), or Cyclophosphamide (CTX, stars, magenta line) as described in Methods. Graphs show (A) percentage of overall survival, (B) percentage progression-free survival, and (C) proteinuria score as measure of clinical efficacy. (D–H) Effect of drug treatment after 62 days at end of study in surviving mice (83 days after AdIFNα induction) on (D) renal glomerulopathy; (E) renal arteritis/periarteritis; (F) splenic peanut agglutinin (PNA+) germinal centers (GCs) (GCs were enumerated in formalin-fixed paraffin embedded [FFPE] spleen sections using IHC); (G) splenic marginal zone B cells (triple immunofluoresence staining of spleen with anti-CD4 (Alexa Fluor 5646, red), anti-Metallophilic Macrophages 1 (MOMA1) (FITC, green), anti-IgM (Aminomethylcoumarin [AMCA], blue) was performed on frozen spleen sections; arrows/+ signs refer to areas of IgM+ marginal zone B cells adjacent to and extending outwardly beyond the MOMA+ macrophage boundary) (20× magnification); and (H) splenic B220+ B cells (B cells were enumerated in FFPE spleen sections using IHC). (I) Serum anti-nuclear antibodies (ANA) Abs were assessed after 7-week treatment for the groups indicated. Statistics indicate (A–C) *P < 0.01, **P < 0.0001, G-744 versus vehicle; *P < 0.05, Cyclophosphamide versus vehicle; *P < 0.05, P505-15 versus vehicle; *P < 0.05, BR3-Fc versus Ig control; ††P < 0.01, G-744 versus BR3-Fc, P505-15, or Cyclophosphamide, respectively. Groups were compared using (A and B) Log-rank Mantel-Cox test or (C) Kruskal Wallis test with Dunn’s correction, (D–F, H, I) one-way ANOVA with Dunnett’s correction or (H) 2-tailed heteroscedastic t tests; symbols represent individual mice.
role of Syk in Akt phosphorylation (Table 2). Conversely, Syk blockade demonstrated a minimal inhibitory effect on Btk-Y223 autophosphorylation. As expected, Syk was essential for PLCγ activation, as P505-15 treatment abrogated BCR-induced Ca2⁺ signaling in a dose-dependent manner (Figure 1E). These data indicate that both Syk and Btk play an essential role in triggering Ca2⁺ flux downstream of the BCR. Interestingly, Syk inhibition had a lesser effect on CD40-mediated murine B cell proliferation (Table 2), indicating that BCR and CD40 signaling pathways may converge at the level of Btk but not Syk in mice.

We next assessed whether specific Btk blockade with G-744 impacts BR3 signaling. G-744 had minimal effect on BAFF-induced B cell survival in vitro and did not alter NFκB p100 processing (Supplemental Figure 1, A and B). Similarly, 1-week in vivo treatment with G-744 did not alter follicular B cell numbers, although it partially reduced splenic marginal zone (MZ) B cell numbers (Supplemental Figure 1, C–E). In contrast, as expected, BR3-Fc treatment significantly reduced both follicular and MZ B cell numbers, while Syk inhibition most significantly lowered newly formed B cell numbers (Supplemental Figure 1, C–E). These data are in agreement with previously published data (35–37) that Btk and Syk are not major constituent of the BR3 survival pathway in follicular B cells and that blockade of BAFF and Btk are likely to have divergent effects on autoimmune responses due to their differential impact on B cell signaling and downstream functions.

As expected from the previously reported role of Syk and Btk in FcR function (18, 21, 33), both Syk and Btk inhibition prevented FcγRIII-triggered TNFα production with similar potency as BCR activation and proliferation (Table 2). These data demonstrate that G-744 is a highly selective Btk inhibitor that prevents both B cell BCR and macrophage FcγRI signaling with similar efficiency. Furthermore, our data indicate that Btk exhibits differential pathway activity in B cells compared with that reported for Syk inhibitors (21, 33).

**Btk inhibition is superior to BAFF blockade or Syk inhibition in preventing IFNα-accelerated lupus and renal pathology.** SLE patients often exhibit higher IFNα levels and type-I IFN activity that are associated with severe disease involving the kidneys, hematopoietic cells, and/or the CNS (38). We examined the efficacy of Btk blockade in preventing LN in the presence of increased levels of IFNα and highly aggressive clinical disease in NZB/W_F1 mice treated with adenovirus-expressing murine IFNα (13, 14) and compared it with BAFF blockade, Syk inhibition, and CTX. Mice treated with Ig control starting 3 weeks after adenovirus expressing IFNα (AdIFNα) injection succumbed to disease 9 weeks after AdIFNα administration. Btk inhibition with 100 mg/kg G-744 significantly improved overall survival (Figure 2A) and progression-free survival (Figure 2B), and stabilized proteinuria (Figure 2C) compared with vehicle. Analyses of drug concentration required to achieve inhibition of pBtk in NZB/W_F1 whole blood in vitro and actual G-744

### Table 1. Potency and selectivity profile of G-744 in enzymatic assays

<table>
<thead>
<tr>
<th>IC₅₀ (nM) at Kₘ&lt;sub&gt;app&lt;/sub&gt;</th>
<th>BTK</th>
<th>BMX</th>
<th>ErbB2</th>
<th>EphA7</th>
<th>FGR</th>
<th>LCK</th>
<th>SLK</th>
<th>SRC</th>
<th>YES1</th>
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<tr>
<td>Selectivity (fold)</td>
<td>1</td>
<td>&gt;2.604</td>
<td>1.367</td>
<td>428</td>
<td>&gt;868</td>
<td>&gt;2.604</td>
<td>&gt;7,813</td>
<td>&gt;2.604</td>
<td>&gt;7,813</td>
</tr>
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</table>

BMX, BM X kinase; ErbB2, epidermal growth factor receptor tyrosine kinase 2 (Her2); EphA7, Ephrin tyrosine kinase receptor A; FGR, Src-family tyrosine kinase Fgr; LCK, lymphocyte-specific tyrosine kinase; SLK, STE20-like protein kinase; SRC, c-sarcoma virus tyrosine kinase; YES1, Yamaguchi sarcoma viral (v-yes) oncogene homolog 1 kinase; Kₘ<sub>app</sub>, apparent Kₘ.

### Table 2. Potency of G-744 and P505-15 in murine and human cell-based assays

<table>
<thead>
<tr>
<th>Stimulus</th>
<th>B cell</th>
<th>Anti-IgM</th>
<th>Anti-IgM</th>
<th>Anti-CD40</th>
<th>BAFF</th>
<th>Anti-FcγRIII</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>pBtk</td>
<td>pAkt</td>
<td>CD86</td>
<td>IL-6</td>
<td>proliferation</td>
</tr>
<tr>
<td>Readout</td>
<td>Mouse</td>
<td>G-744</td>
<td>20</td>
<td>&gt;10000</td>
<td>64</td>
<td>54</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P505-15</td>
<td>784</td>
<td>169</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td></td>
<td>Human</td>
<td>G-744</td>
<td>15</td>
<td>&lt;50%</td>
<td>nd</td>
<td>28</td>
</tr>
</tbody>
</table>

Values given are mean of the IC₅₀ values (nM) for B cell and macrophage assays with the indicated stimulation conditions; nd, not determined.
plasma drug concentrations from the animals in the pharmacokinetic study group demonstrated that drug levels that were at least 70% inhibitory for pBtk were maintained throughout the dosing regimen (Supplementary Figure 2, A and C). In comparison, a similar extent of Syk inhibition at 60 mg/kg P505-15 (based on P505-15 concentration required to suppress FcεRI-induced CD63 levels on basophils in whole blood, Supplemental Figure 2B and ref. 32) significantly improved progression-free survival (Figure 2B) but only showed a trend of ameliorating overall survival compared with vehicle ($P = 0.2$; Figure 2A) and did not prevent progression of proteinuria. BR3-Fc showed a significant effect on overall survival but not on progression-free survival ($P = 0.4$) or proteinuria. CTX improved overall survival, progression-free survival, and proteinuria. Interestingly, Btk inhibition at 100 mg/kg showed markedly increased overall survival, compared with BR3-Fc, and progression-free survival, compared with all 3 alternate treatments (P505-15, BR3-Fc, and CTX). Furthermore, G-744 was superior in preventing proteinuria compared with P505-15 and BR3-Fc. At the end of the study, G-744 showed a trend in improving severity of glomerulopathy and significantly suppressed arteritis/periarteritis compared with surviving mice in the vehicle group with a relative effect at least comparable with that of CTX. Of note — in an alternate IFNα-accelerated lupus study in which NZB/W_F1 mice were treated for only 4 weeks with G-744, BR3-Fc, or CTX to allow for a greater number of surviving mice in the control groups for comparison — G-744 treatment significantly prevented both glomerulopathy and arteritis/periarteritis compared with vehicle and glomerulopathy compared with BR3-Fc (Supplemental Figure 3, A–D). Syk inhibition did not significantly improve renal histopathology parameters (Figure 2, D and E).

Btk inhibition reduces GCs and Ab production in both IFNα-accelerated and spontaneous lupus. To assess the effects of sustained drug treatment on splenic B cell populations, we examined spleen sections in surviving mice from the IFNα-accelerated lupus study using IHC or immunofluorescence. Treatment with G-744 significantly decreased GC frequency by 60% compared with vehicle, while BR3-Fc, CTX, and P505-15 exhibited only a minimal effect in suppressing splenic GC formation (Figure 2F) and G-744 treatment was superior in preventing MZ B cell loss (Figure 2G), with BR3-Fc, P505-15, or CTX having no discernible effect on splenic MZ morphology. BR3-Fc treatment also decreased the number of splenic B220+ B cells as deter-

Figure 3. Effect of G-744 on splenic B, T, and plasma cells in IFNα-accelerated NZB/W_F1 lupus model. Fourteen-week-old preautoimmune NZB/W_F1 female mice (n = 10/group) were administered with adenovirus containing murine IFNα. Drug treatment was started 3 weeks after AdIFNα injection with 100 mg/kg G-744, vehicle HPMC, 5 mg/kg BR3-Fc, 5 mg/kg Ig control Abs, or Cyclophosphamide (CTX) as described in Methods and continued for 4 weeks. Graphs show the number and phenotype of splenic lymphocyte population from individual surviving mice as determined by flow cytometry and expressed as mean ± SEM, symbols represent individual mice: (A) number of proliferating B cells (CD5–B220+Ki67+); (B) number of B2 cells (B220+CD38+); (C) CD86 levels on B cells (CD5–B220+); (D) ICOS levels on CD4 T cells (CD5+CD4+); (E) number of GC B cells (B220+CD38+); number of (F) total, (G) IgM, (H) IgG1, (I) IgG2a, and (J) IgG2b splenic CD138−B220− PCs. Significance indicate *$P < 0.05$, G-744 or CTX versus vehicle, BR3-Fc versus Ig control (one-way ANOVA with Dunnett’s multiple comparison test); $\wedge P < 0.05$, compare G-744 with BR3-Fc (2-tailed Student’s t test with Welch’s correction).
Figure 4. Gene expression changes in spleen after cyclophosphamide treatment or blockade of Btk, Syk, or BAFF in IFNα-precipitated LN in NZB/W_F1 mice. LN was IFNα-accelerated in female NZB/W_F1 mice as described in Figure 3. Mice were treated for 4 weeks starting 3 weeks after AdIFNα induction with vehicle, 100 mg/kg G-744, 60 mg/kg P505-15, 7.5 mg/kg Ig control, 7.5 mg/kg BR3-Fc, or Cyclophosphamide (Cytoxan/CTX) (n = 10/group), followed by RNA sequencing analysis of splenic RNA. Two groups of 17-week-old NZB/W_F1 animals were also included (see Supplemental Figure 7) that were either not injected with AdIFNα (naive group) or treated for 3 weeks with AdIFNα (AdIFNα group). (A) Gene set enrichment analysis was performed using predefined immune gene modules (n = 5/group). Gene sets that showed significant enrichment (permutation \( P < 0.01 \)) are shown as a heatmap indicating the magnitude of the gene set enrichment statistic. (B–D) G-744 reduces transcripts associated with ASC generation. (B) Calculated fold-changes in spleen from vehicle-treated versus naive animals (x-axis), or G-744-treated versus vehicle-treated animals (y-axis), are shown for all transcripts (gray) or ASC-specific transcripts (black). Dotted lines indicate a 2-fold change in expression. (C) Heatmap visualization of select ASC-specific transcripts that were induced (log2-fold change > 2, \( P < 0.05 \)) in spleens of vehicle-treated animals compared with naive animals. Each column in the heatmap represents 1 animal within each treatment group. Genes marked in blue were reduced after G-744 treatment compared with
vehicle-treated animals (fold change > 1.5, P < 0.05). (D and E) Box plot of reads per kilobase of transcript per million mapped reads (RPKM) expression of (D) select individual genes from C significantly affected by G-744 treatment and (E) select B cell genes affected by BR3-Fc treatment. Gene expression data were plotted using box plots overlaid with all observed data points. The box corresponds to the 1st quartile, median (black line), and 3rd quartile. The whiskers extend to the most extreme data point that is no more than 1.5 times the interquartile range from the box; *P < 0.05, **P < 0.005, ***P < 0.0005, using DESeq2 to fit a negative binomial model, as described above.

mined by evaluating the B220+ immunoreactivity in spleen sections (Figure 2H). These findings are consistent with the role of BAFF in B cell survival. In comparison, CTX and P505-15 treatment showed a modest effect and G-744 a minimal effect on B220+ reactivity. Examination of serum autoantibodies at 7-week treatment showed that ANA Abs were significantly reduced by G-744, but not by the other treatments (Figure 2I). Anti-dsDNA Abs showed a trend of reduction by G-744 for IgM isotypes (Supplemental Figure 4A) but no significant effect for IgG3-specific anti-dsDNA Abs (Supplemental Figure 4B). In comparison, BR3-Fc significantly reduced IgG3 anti-dsDNA Abs. Similar results were obtained in an alternate study where we compared Btk inhibition at 35 mg/kg or 100 mg/kg of G-744 to that of BAFF blockade or CTX treatment in IFNα-accelerated NZB/W_F1 lupus (Supplemental Figure 5, A–G). We further analyzed total serum Ig concentrations in this model and observed that G-744 reduced titer of IgM and IgG3 Abs (Supplemental Figure 4, C and F) but did not affect titers of IgG1 or IgG2a (Supplemental Figure 4, D and E).

Given previous reports that Btk inhibitor treatment in spontaneous lupus disease in NZB/W_F1 mice without IFNα-acceleration reduced IgG dsDNA Ab titers (23, 24), we also examined the effect of highly selective Btk inhibition with G-744 on disease parameters and autoantibody titers in NZB/W_F1 mice with spontaneously occurring disease and compared it to BR3-Fc and CTX. Therapeutic treatment with G-744 at 35 mg/kg and 100 mg/kg in contrast to BR3-Fc significantly improved overall survival (Supplemental Figure 6A) and progression-free survival (Supplemental Figure 6B), and it stabilized proteinuria (Supplemental Figure 6C); it also showed a significant dose-responsive inhibitory effect of 100 mg/kg G-744 treatment on GC formation compared with CTX (Supplemental Figure 6D). Surprisingly, in contrast to previously published data, BR3-Fc in this study did not prolong survival, perhaps due to the late treatment start at 36 weeks of age and the highly advanced proteinuria in this cohort. G-744 at 100 mg/kg, but not BR3-Fc or CTX, significantly reduced serum anti-dsDNA (Supplemental Figure 6E) and anti-histone (Supplemental Figure 6F) autoantibody levels after treatment for 56 or 112 days. Furthermore, gene expression analyses from whole blood demonstrated that expression of B cell (Ighg1), myeloid cell (Csf1r), and IFN-regulated genes (Ir7, Oas3, Cmpk2, Mx1) were significantly decreased over time with G-744 treatment (Supplemental Figure 6G).

Taken together, these results indicate that highly selective Btk inhibition is superior to BAFF blockade, Syk inhibition, and CTX in ameliorating disease parameters and autoantibody production in the IFNα-accelerated and spontaneous NZB/W_F1 lupus models.

G-744 significantly reduces splenic PCs. Given the effect of selective Btk antagonism on GC frequency and autoantibody production in therapeutic treatment regimens, we examined in more detail the mechanism of action of Btk inhibition with G-744 treatment on B cell subsets in spleen in the presence of elevated IFNα levels. Btk inhibition caused a potent decrease in the number of proliferating Ki67+ B cells compared with vehicle, BR3-Fc, and CTX-treated mice after 4 weeks of treatment (Figure 3A). Consistent with the effect on proliferating B cells, G-744 but not BR3-Fc reduced the number of total B2 cells (Figure 3B). Antagonism of Btk, but not the other treatments, decreased cell CD86 and CD4+ T cell ICOS levels (Figure 3, C and D), as well as the number of GC B cells (Figure 3E) compared with control-treated mice. Importantly, G-744 significantly diminished the total number of splenic PCs compared with BR3-Fc and vehicle (Figure 3F). Furthermore, Btk inhibition reduced the number of splenic IgM+ plasmablasts, as well as all isotype-switched PCs tested (IgG1+, IgG2a+, and IgG2b+) (Figure 3, G and H). In comparison, BR3-Fc did not significantly lower the numbers of total, IgM+, or isotype-switched PCs compared with control Ig-treated mice. CTX moderately lowered the number of total, IgG2a+, and IgG2b+ PCs and significantly reduced IgG1+ PC in the spleen (Figure 3, F and G).

G-744 inhibits splenic gene expression associated with B cell terminal differentiation. To investigate potential mechanisms by which above treatments influence LN and tissue homeostasis during disease progression, we performed transcriptional profiling by RNA sequencing of whole spleen RNA isolated from AdIFNα-injected NZB/W_F1 mice after 4-week treatment with G-744, P505-15, BR3-Fc, CTX, or controls. Drug treatment started 3 weeks after AdIFNα injection and onset of proteinuria when...
mice were 17 weeks old (proteinuria ≤ 100 mg/dl, Supplemental Figure 7A). AdIFNα-treated 21-week-old NZB/W_F1 mice in the vehicle or Ig control group at the end of the study were nephritic (at least 80% of mice with proteinuria ≥ 300 mg/dl), whereas all drug treatment partially prevented progression of proteinuria. For comparison, we included a group of 17-week-old NZB/W_F1 animals that were not injected with AdIFNα showing no or minimal proteinuria (termed naive group) and a group of 17-week-old mice that were terminated after 3-week AdIFNα induction (AdIFNα group).

Gene set enrichment analysis (GSEA) of spleen using gene sets describing antibody-secreting cells (ASC), other immune cells, inflammation, and general repertoire modules (39–41) demonstrated that G-744 treatment decreased gene expression associated with inflammation and cell death in the spleen (Figure 4A). Consistent with our in vivo findings, there was a strong inhibitory effect of G-744 treatment on modular repertoires associated with immune cell types including B cells, PCs, myeloid cells, and monocytes in the spleen (Figure 4A). Of particular note, the gene set that had the strongest inhibition due to Btk inhibition was the ASC set (Figure 4, A–D) that was recently defined (41). In comparison, other treatments did not result in decreases of equal magnitude. BR3-Fc treatment had moderate effects on select predefined B cell and cell cycle modules, but not ASCs. Specifically, BR3-Fc downmodulated expression of B cell lineage marker genes in the “B cell M4.10 module” including Cd19, Cd79a, Blk, Bank1, and Ms4a1 (Cd20), in accordance with its role in B cell survival (Figure 4E). Few changes in gene expression were observed in CTX or Syk inhibitor treatment, except upregulation of a neutrophil gene set by CTX and a mild decrease in ASC-associated gene expression after Syk inhibition (Figure 4A).

Examination of changes in individual ASC genes revealed that the majority of these genes were induced in animals with disease as compared with naive controls (Figure 4B, indicated by the back dots), and expression of these same genes was downmodulated by G-744 treatment. Further analyses of these ASC-specific transcripts that were induced (fold change > 2, \( P < 0.05 \)) in vehicle-treated compared with naive animals showed that G-744 normalized expression of the majority of them, while Syk inhibition, BAFF blockade, and CTX exhibited little effect (Figure 4C). These genes, including Sec24a, Edem1, Fkbp11, and Xbp1 (Figure 4, C and D), encompassed biological function connected with Ab secretion such as the unfolded protein response, protein post-translational modifications, folding, and transport in the ER.
Figure 6. Gene expression changes in kidney in IFNα-accelerated LN in NZB/W F1 mice after treatment with G-744, P505-15, BR3-Fc, or cyclophosphamide (Cytoxan). RNA sequencing analysis was performed on kidney RNA (n = 5/group) derived from the experiment described in Figure 4. (A) Gene set
enrichment analysis of predefined immune (see Figure 4A) and human LN ortholog gene modules (as defined in D and E); gene sets that showed significant enrichment (permutation \( P < 0.01 \)) are shown as a heatmap indicating the magnitude of the gene set enrichment statistic. (B and C) Analyses of overlapping effects on renal gene transcription by drug treatments. (B) Renal transcripts that were significantly upregulated in diseased kidneys versus naive animals (2-fold elevated with FDR of 0.05) were assessed for their downregulation (±1.5-fold with FDR of 0.05) by cyclophosphamide (CTX) versus G-744 treatment. Blue dots indicate genes significantly reduced by G-744, black dots indicate genes significantly reduced by CTX, and orange dots indicate genes significantly downregulated by both treatments. (C) Venn diagram showing the degree of overlap of genes upregulated in the kidneys of diseased animals that were subsequently decreased by CTX, G-744, or P505-15 treatment using the same statistical cutoffs as stated for B. The number of genes affected by each treatment is indicated within each intersection, and the number of genes unaffected by any of these treatments is indicated outside of the circles. (D and E) Effect of drug treatment on human LN ortholog genes previously identified to be upregulated in the (D) glomerulus or (E) tubulointerstitium of nephritic kidneys from human LN patients (see Methods). Genes that showed individual significant changes with any treatment versus their controls are indicated (±1.5-fold difference between groups with an adjusted \( P < 0.01 \) as determined using a negative binomial generalized linear model). Each column in the heatmap represents 1 animal within each treatment group.

G-744 treatment reduces IgM and IgG deposition in kidney glomeruli and proximal convoluted tubules. In light of the profound effects on ASC gene expression in the spleen of Btk inhibitor–treated NZB/W_F1 mice, we assessed the effect of specific Btk inhibition compared with Syk blockade and CTX on renal IgM and IgG deposition from the experiment described in Figure 4. Proteinuric vehicle-treated animals showed an increase in IgM staining in glomeruli (Figure 5A), and quantification of these deposits revealed a statistically significant increase in IgM-rich inclusions in both glomeruli and proximal convoluted tubules in diseased versus naive mice (Figure 5, B and C). In contrast, there was a much less pronounced increase in glomerular IgG deposition with disease progression compared with glomerular IgM and tubular IgG deposits (Figure 5, B, D, and E). G-744 treatment resulted in a significant reduction in both glomerular and tubular IgM deposits (Figure 5, B and C), while IgG deposits were significantly reduced only in the tubules (Figure 5, D and E). P505-15 treatment significantly reduced tubular but not glomerular IgM scores, whereas CTX reduced glomerular but not tubular IgM scores (Figure 5, B and C). Significant but lower magnitude decreases in tubular IgG scores were also observed in P505-15– and CTX-treated animals. In comparison, BR3-Fc treatment showed no significant decrease in tubular scores (Figure 5E). Further examination of IgG in the glomeruli showed that none of the drugs after 4 weeks of treatment significantly lowered glomerular IgG deposits (Figure 5D), although glomerulopathy was significantly reduced by G-744 treatment (Supplemental Figure 7B) as previously observed (Supplemental Figure 3B).

Btk inhibition and CTX treatment show overlapping effects in reducing renal gene expression linked to LN in NZB/W_F1 mice. To determine effects of drug treatment on LN in mouse kidneys, we performed transcriptomic profiling of whole kidney RNA from the study in Figure 4 using RNA sequencing coupled with GSEA analyses using inflammatory and general repertoire modules examined above for spleen (Figure 6A). G-744 treatment showed a pronounced effect on gene modules reflective of myeloid cell activation, cell cycle, and several inflammation repertoires, while Syk inhibition reduced gene transcription associated with only one of the inflammation modules. BAFF blockade had little effect on global renal gene transcription. CTX treatment decreased transcriptional modules linked to protein synthesis, select inflammatory patterns, and immune responses associated with antigen presentation — in particular H2, class-II MHC gene expression. Of note, CTX increased erythrocyte transcriptomes and displayed mixed effects on cell cycle modules that were either under- or overexpressed (Figure 6A).

Analyses of genes increased during IFNα-accelerated nephritis and downregulated by more than one drug treatment revealed that Btk inhibition and CTX treatment demonstrated overlapping effects on decreasing disease-induced genes: of 1,131 upregulated transcripts (>2-fold induced in vehicle versus naive group, with an adjusted \( P < 0.05 \)), 46% of the transcripts (518 total) were downregulated by both G-744 or CTX treatment at least 1.5-fold, with an adjusted \( P < 0.05 \) (Figure 6, B and C). In comparison, Syk inhibition only reduced a subset of the transcripts concomitantly decreased by G-744 or CTX (45%, 234 total) and showed minimal unique overlap with CTX (H2 class-II MHC, CD74) or G-744 (PTPRN) (Figure 6C).

The most highly expressed genes during onset of clinical nephritis that were reduced either by G-744 or CTX were known mediators of tubulointerstitial damage or fibrosis (IL1-F6, Grem1, Lcn2) and tissue remodeling associated with myeloid cell recruitment and activation (Cxcl17, Timp1). Unique nephritis-induced genes downregulated predominantly by Btk inhibition included myeloid cell receptor genes (Cec4d [MCL], Clec4n [Dectin-2], Itgam, Trem1), genes associated with B cell proliferation (Fcr15), and several transcription factor–encoding genes (Btg2, Egr1, Fos, Fosb, Irf4, Jun).

Btk inhibition reduces renal gene expression associated with LN in human patients. To understand the possible relevance to human disease, we intersected genes previously shown to be elevated during nephritis in
kidneys isolated from different lupus mouse strains with genes upregulated in (i) the glomeruli, (ii) the tubulointerstitium, or (iii) both compartments from kidney biopsies of human LN patients versus normal control subjects (42) and assigned them to 3 modules (Figure 6A). Examination of drug treatment effects on gene expression associated with human LN in these renal compartments demonstrated that therapeutic treatment with G-744 significantly abrogated transcripts associated with both glomerular and tubular gene expression (Figure 6, D and E; and Supplemental Figure 8A). CTX treatment significantly inhibited the transcriptome of the tubulointerstitial compartment with a lesser effect on the glomerular module compared with G-744. In comparison with G-744 and CTX, BR3-Fc or P505-15 showed a minimal effect (Figure 6, D and E; and Supplemental Figure 8A). Assessment of individual genes defined by Berthier et al. for the glomerulus and the tubulointerstitial compartment (Figure 6, D and E) or genes common to both compartments (Supplemental Figure 8A) confirmed that the genes were elevated in animals with active disease (vehicle and Ig drug-control groups) versus animals with minimal disease (naive group) (42). Interestingly, while the AdIFNα group showed a similar extent of ASC-associated transcript induction compared with vehicle/Ig control groups (Figure 4C), nephritis-induced gene expressions was not fully developed in the AdIFNα group compared with control groups, in agreement with relative disease activity exemplified by the higher proteinuria and glomerulopathy scores in the control groups (Supplemental Figure 7). Btk inhibitor–regulated genes expressed in the glomerulus only or in both compartments were enriched for genes encoding molecules associated with myeloid cells and inflammation (Cd14, Fgr1, Fpr1, IgF6f, IL1Rn, Msrl, Plaur, Spltc2, Tlr1), cell cycle function (Ccnb2, Cdc20, Ckap2, Fam57a, Fam129a, Gins2, Nusap1, Prc1, Rad51ap1, Top2a), cell migration or adhesion (Ccr2, Cd44, Cxcr4, Hmr [Cd168], Itgam [Cd11b], Lgals3, Rgs1), and extracellular matrix function and degradation (Adamts1, Mmp2, Col3a1, Timp1, Fn1) (Supplemental Figure 8, B and C).

Genes exclusively associated with tubule expression and inhibited by G-744 were enriched for additional molecules associated with extracellular matrix function and adhesion regulation (Cdh3, Col1a1, CXCL10, Gpr126, Tpg, Vcam1), as well as molecules that control membrane trafficking and may be associated with cell–extracellular matrix (ECM) interactions (Anxa1, Anxa2, Golm1, Kdelr3, Myof, Ppl2, Splt) (Supplemental Figure 8D). Several genes with less well-characterized functions — but with potential roles in cell proliferation, cell death, angiogenesis, or vascular function (Ankr6d, Anxa3, Asns, F2r1l, Krt19, Tmsb10, Tsc22d1, Tuba1a, Ywha [14-3-3η]) — were also modified by Btk inhibition.

Therefore, short-term and highly specific inhibition of Btk in the NZB/W_F1 model affected both B cell and myeloid cell responses in the secondary lymphoid system and further decreased the expression of
renal ortholog genes previously identified to be differentially expressed in the glomerular and tubulointerstitial compartments in human LN patients.

**G-744 inhibits human B cell activation and differentiation into plasmablasts.** To examine whether effects observed with Btk inhibition in the mouse system translate to human, we assessed Btk pathway activity in human B cells and monocytes. In response to IgM activation, G-744 blocked pBtk with an IC₅₀ of 15 nM (Table 2), while pAkt phosphorylation was not significantly changed. Furthermore, BCR-induced B cell proliferation and IL-6 production was inhibited with similar potency. B cell proliferation was also potently abrogated after CD40L stimulation. In comparison, BAFF-induced B cell survival was only partially reduced at G-744 concentrations that blocked BCR-triggered responses, albeit with a relatively higher impact of Btk inhibition on BAFF-mediated B cell survival in human B cells as compared with mouse B cells. As observed in murine B cells (Supplemental Figure 1B), Btk inhibition did not affect NFκB p100 to p52 processing (data not shown). Finally, G-744 prevented FcγRIII-mediated TNFα production in human monocytes (Table 2).

Given the reduction of PC generation by G-744 in NZB/W_F1 LN, we explored whether Btk contributes to TLR9-induced human memory B cell differentiation into plasmablasts. Stimulation of CD27⁺ memory B cells with CD40L or TLR9 ligands in the presence of cytokines including IFNα (43) yielded about 50% or 15% plasmablast (CD20⁻CD38⁺⁺), respectively (Figure 7, A and B). G-744 potently abrogated both CD40L- (Figure 7, A and B) and ODN2006-induced (Figure 7, B and D) plasmablast generation in a dose-dependent manner. Importantly, we also observed that, while Btk is expressed in differentiated plasmablasts at a level comparable with naive B cells, inhibition with G-744 did not affect survival of plasmablasts and PCs in vitro (Supplemental Figure 9). These data demonstrate that Btk activity is essential for B cell proliferation and differentiation into plasmablasts in response to either CD40 or TLR9 activation in the presence of IFNα, but it does not appear to be required for survival of these cells once they have differentiated.

**Discussion**

LN is a major cause of morbidity in SLE patients for whom no immune-based targeted therapies are currently approved (15, 16). We developed an oral highly specific reversible Btk inhibitor, G-744, with over 400-fold selectivity against all 284 kinases tested, including highest selectivity against Tec- and Src-family kinases compared with other Btk inhibitors (Supplemental Table 1). Specific Btk inhibition shows a profound effect on improving survival, as well as preventing glomerulonephritis and progression of proteinuria in animals with severe disease when high levels of IFNα are present in the IFNα-accelerated lupus disease (Figure 2) or the spontaneous NZB/W_F1 lupus model (Supplemental Figure 6). Notably, G-744 treatment is the most efficacious from all treatments tested and superior to CTX (15, 16), the BAFF antagonist BR3-Fc (17), and the Syk inhibitor P505-15 (32). Importantly, in NZB/W_F1 mice with IFNα-accelerated lupus disease, our results demonstrate a global effect of specific Btk blockade on downmodulating both the transcriptome associated with PC generation in the spleen, in particular secretory pathway genes, as well as renal gene expression associated with LN in human SLE patients.

Interestingly, CTX and G-744 showed overlapping effects on inhibiting renal gene transcription associated with human LN, while CTX did not affect the PC program. BR3-Fc treatment, although moderately prolonging survival of mice, did not impact gene transcription associated with PC generation or nephritis, which is consistent with the modest disease-modifying effects observed in SLE patients. Surprisingly, Syk inhibition did not match Btk inhibition, and while P505-15 showed a similar trend in improving disease parameters compared with G-744 at drug levels that lead to comparable respective enzyme inhibition (Supplemental Figure 2), P505-15 treatment only minimally reduced gene transcription controlling B cell terminal differentiation and tubular damage. These results identify Btk as a gatekeeper of processes controlling glomerulonephritis in NZB/W_F1 mice with high levels of IFNα and provide insights that Btk-dependent pathways that drive disease in mice may also be active in human SLE.

IFNα triggers pleotropic effects on the immune system in both human and mice, including sustained B cell proliferation, upregulation of TLR7, activation of DCs and myeloid cells, induction of T-dependent immune responses, and impairment of GC selection resulting in increased autoreactivity of the antigen-driven B cell repertoire (7, 8, 10, 11, 14). AdIFNα promotes a feed-forward amplification process in lupus-prone NZB/W_F1 mice triggering lupus disease that exhibits distinct features compared with the spontaneous model, including requirement of T cells for initiating disease, persistent production of short-lived PCs, enhanced
expression of Btk triggers increased PC numbers, anti-nucleosome autoantibody production, and SLE-like disease-promoting effects of IFNα in both human and mice, we chose to analyze the consequences of selective Btk inhibition in the AdIFNα-accelerated NZB/W_F1 lupus model in particular, as this model is more resistant to therapeutic intervention, including CTX in combination with anti-CD40L and CTLA4-Ig (14).

Btk inhibition for 4 weeks at drug levels of G-744 that inhibit Btk enzyme activity throughout the dosing period ablated in the spleen the number of proliferating B cells, activated CD86+ B cells, GC B cells, and PCs (including IgM+, IgG1+, IgG2a+, and IgG2b+ PCs), as well as ICOS+ CD4 T cells of AdIFNα-treated NZB/W_F1 mice (Figure 3). In contrast, BR3-Fc treatment did not significantly affect splenic GC B cell or above IgG+ PC populations, but it did show a trend in reducing proliferating B and B2 cells, while CTX had a modest effect on IgG2a+ and IgG2b+ PCs with little effect on B cell subsets. PCs were also reduced in the BM by Btk inhibition (data not shown), although —consistent with previous data — only low numbers of PCs were detected in the BM in vehicle-treated mice, in favor of large numbers of short-lived PCs in the spleen (14). Given the extensive effect on PC numbers, it is surprising that, while we did detect a significant reduction in ANA autoantibodies (including IgM, IgG, and IgA isotypes) measuring reactivity to various antigens (Sm, RNP, SSA/Ro, SSB/La, DNA, Scl70, and Jo-1), we did not observe a significant effect of G-744 after 6 weeks of treatment on anti-dsDNA IgG or IgG3 autoantibody titers in the presence of AdIFNα (Supplemental Figure 5E and Supplemental Figure 4B). However, anti-dsDNA IgM autoantibodies showed a trend of reduction ($P = 0.077$), and total IgM and IgG3 serum Ab concentrations were significantly reduced after 6 weeks of treatment with G-744 (Supplemental Figure 4 C–F), indicating a potent effect of G-744 treatment (i) on IgM production by short-lived plasmablasts that are predominant in this model or (ii) on alternate ANA Ab species. In comparison, anti-dsDNA and anti-histone Ab titers were significantly reduced with 100 mg/kg G-744 in the spontaneous model after 6 weeks of treatment, in agreement with effects observed with less selective and/or irreversible Btk inhibitors in the spontaneous N XB/W_F1 model (23, 24) and in the BXSB-Yaa or the pristane DBA/1 lupus model that have a greater contribution of TLR7 and TLR7/type-I IFN pathways, respectively (29). Although unclear at present, the lack of G-744 effect in the IFNα-accelerated model, specifically on dsDNA IgG autoantibody titers that are particularly dependent on IFNα/TLR9 signaling, may be due to the continued expression of high levels of exogenous adenoviral-derived IFNα.

In this respect, Btk inhibition reduced the peripheral blood IRG signature that accompanied severe disease onset around 36 weeks in the spontaneous model (Supplemental Figure 6G) but did not significantly impact the blood IRG signature in the IFNα-accelerated model (data not shown). Priming B cells with IFNα lowers the BCR activation threshold and may trigger spontaneous peripheral activation of anti-DNA B cells in a TLR9-dependent manner and their differentiation into Ab-forming cells via an extrafollicular pathway (45, 46), which may be less dependent on BCR/Btk signaling and T cell help.

To further understand the mode of action of selective Btk inhibition in reducing PCs and GCs in spleen, genome-wide transcriptional profiling followed by GSEA identified a predominant role of Btk in controlling gene transcription associated with B cell terminal differentiation and ASCs, in addition to strong inhibitory effects on myeloid cells and monocyte modules. Btk inhibition resulted in downregulation of genes involved in most stages of the secretory process encoding molecules associated with folding of ER proteins, including ER chaperones (Fkbp11, Txnndc11, Txndc5), degradation of misfolded ER proteins by the ERAD pathway (Edem1, -2, -3), protein glycosylation (Man1a, Manea, Fut8), and ER-Golgi vesicular trafficking (Sec24d, Sec24a). The main transcription factor that controls the secretory capacity of PCs is XBP1, and transcription of most of the above secretory pathway genes has been shown to be dependent on XBP1 (47). Accordingly, splenic XBP1 expression was also strongly inhibited by Btk inhibition and so were its regulators Ern1/IRE1 and Prdm1/Blimp1, as well as additional Blimp1-controlled genes (Sdc1, ELL2, IgJ) essential for PC terminal differentiation. Furthermore, G-744 inhibited IRF4 expression that, at higher doses, controls the initiation of PC differentiation. Downregulation of select splenic genes related to B cell differentiation, in particular Blimp1, have previously been noted in the spontaneous NZB/W_F1 model with RN486 (24), indicating that the transcriptional program related to B cell terminal differentiation may not be highly controlled by IFNα. Unlike Btk inhibition, BAFF blockade had no effect on ASCs but had moderate effect on B cell marker- and cell cycle–associated gene expression, demonstrating that Btk inhibition and BR3-Fc exhibit divergent effects on B cell homeostasis. These data reveal Btk as a potential candidate to control an essential checkpoint in lupus B cell pathogenesis in the presence of IFNα, as Btk blockade ablated most aspects of the PC program in NZB/W_F1 mice. In this respect, forced overexpression of Btk triggers increased PC numbers, anti-nucleosome autoantibody production, and SLE-like
Furthermore, suppression of PC generation may translate to humans, as Btk inhibition with G-744 prevented in vitro differentiation of human memory B cells to plasmablasts in response to CD40L or TLR9 ligands plus IFNα.

The molecular mechanisms of how Btk regulates the PC program during terminal B cell differentiation are currently unknown, but Btk may exert its function due to its prominent role in Ca²⁺ homeostasis, controlling inositol trisphosphate generation and the depletion of thapsigargin-sensitive intracellular calcium stores in the ER, as well as sustained extracellular Ca²⁺ influx (49). Pronounced alterations in B cell activation thresholds due to signaling abnormalities and continued stimulation with autoantigens particularly after a break in B cell tolerance may lead to sustained BCR activation, as well as prolonged depletion of ER Ca²⁺ stores and extracellular Ca²⁺ influx in a Btk-dependent manner. Accordingly, peak BCR-triggered [Ca²⁺]i responses were shown to be significantly higher in B cells from lupus patients compared with normal controls (50).

However, changes in BCR and Ca²⁺ signaling are likely only one component of many altering loss of B cell checkpoint function. Thus, deregulated CD40 pathway activity has been implicated in promoting pathogenesis in SLE (51), and Btk inhibition blocked anti-CD40–mediated B cell proliferation in murine and human B cells. In this respect, it is relevant to note that autoimmune pathology in CD19-hBtk mice is dependent on B-T cell interaction through CD40L and CD19-hBtk B cell–promoted ICOS T cell expression (52), consistent with our data demonstrating suppression of ICOS expression on CD4 T cells (Figure 3D). Furthermore, stimulation of B cells with anti-CD40 (or anti-IgM, LPS, or CpG) upregulates Btk protein expression (48). Interestingly, Arid3a/Bright is induced in mature resting B cells after BCR and CD40 activation (53), and Arid3a was among the more highly Btk-dependent transcripts downregulated in lupus spleen. The low magnitude of prevention from anti-CD40–triggered B cell proliferation by Syk antagonism, which has been previously observed (37), may provide one explanation why P505-15 is less effective than G-744 in reducing B cell–centered responses. Other mechanisms may include the minimal effect of Syk inhibition preventing Btk activation and the more potent antagonism by P505-15 of Akt compared with G-744.

In addition to pathogenic functions in LN directly mediated by B cells, IC deposition accompanied by macrophage and DC infiltration is a hallmark of inflammatory glomerulonephritis in SLE patients and correlates with poor renal outcome (4). Macrophages play a central role in driving inflammation through recognition of IC via TLRs and FcγRs, resulting in effector functions including production of key inflammatory cytokines such as IL1β, IL-18, and TNFα. In the IFNα-accelerated NZB/W_F1 lupus model, glomerulonephritis is characterized by increased proliferative lesions with de novo crescent formation and significantly higher numbers of infiltrating CD11b+Gr1−CD11c+ F4/80+ macrophages than in spontaneously nephritic NZB/W_F1 mice, and these pathogenic findings are suppressed by deletion of macrophages using clodronate (5). We demonstrated previously the crucial role of Btk in controlling IC-mediated arthritis (18). According to these findings, G-744 inhibited FcγR-triggered TNFα production in mouse macrophages and human monocytes (Table 2), as well as in myeloid cell–dependent CIA in Lewis rat (data not shown). Although not formally shown in the IFNα-accelerated model, activating FcRs on circulating hematopoietic cells are necessary for IC-controlled renal pathogenesis in NZB/W_F1 (6). Moreover, activation of FcγR–CD11b+ myeloid cells, including macrophages, by glomerular IC deposits is sufficient to initiate glomerulonephritis.

To evaluate the impact of Btk inhibitors on myeloid cell–dependent glomerulonephritis and identify potential Btk-dependent pathway activity in human renal pathogenesis, we characterized the functional consequences of specific Btk inhibition on transcriptional modules that drive renal disease in mice and that also overlapped with characterized renal gene expression patterns of human lupus patients (42). We observed major changes in transcription after treatment with CTX or G-744, likely reflecting amelioration of disease. Surprisingly, Btk inhibition demonstrated a significant overlap in transcriptional response with CTX of 66% and 48% in tubulointerstitial and common glomerular/tubulointerstitial gene sets, respectively. Further, G-744 and CTX treatment had overlapping effects in decreasing expression of about 46% of gene orthologs that were previously identified as being elevated in the kidneys of LN patients. Therapeutic Btk inhibition most significantly abrogated transcripts associated with renal inflammation, cell migration, and myeloid cell infiltration of the glomerulus, but also showed a major effect on tubular gene expression — in particular on molecules associated with membrane trafficking and function that may communicate with the extracellular matrix. These data are consistent with changes in renal gene expression by inhibition of Btk in a nephrotoxic serum transfer model of glomerulonephritis (30).
Accompanying the gene expression changes was a significant decrease in IgM deposition in glomeruli as well as IgM and IgG staining in proximal convoluted tubules after 4-week inhibitor treatment. In comparison, we did not observe decreased IgG deposition in the glomeruli at this time point, and glomerular IgG deposition was only moderately increased with disease induction. While we cannot exclude that isotype-specific or select tissue antigen–directed glomerular IgG deposits were missed in our studies due to availability of only formalin-fixed tissue for analyses, the more profound increase in glomerular IgM deposits with disease and concomitant G-744 treatment effects is consistent with its stronger effect on IgM production. Nevertheless, the significant reduction of IgG deposition in proximal convoluted tubules may have contributed to decreased activation of tubular epithelial cells and tubulointerstitial pathology. In this respect, approximately 70% of LN patients have demonstrable immune aggregates along the renal tubular basement membrane, and the severity of tubulointerstitial changes inversely correlate with renal prognosis (54).

In conclusion, selective inhibition of Btk results in superior efficacy in spontaneous and IFNα-accelerated NZB/W_F1 lupus models, compared with BR3-Fc, CTX, and Syk inhibition, by regulating expression of genes that are involved in PC differentiation and Ab secretion, as well as genes involved in glomerular and tubulointerstitial renal pathology during LN in human patients.

Methods

Btk enzymatic activity assays and selectivity screens. The in vitro biochemical assay quantified Btk-catalyzed tyrosine phosphorylation of a synthetic peptide, and the kinome selectivity of G-744 was tested in duplicate at 1 μM against a panel of 285 active recombinant human kinases. Further details are given in Supplemental Methods.

Drug formulation and dosing. In vitro studies were performed using crystalline compound dissolved in DMSO. Prior to activation, cells were incubated for 30 min to 1 h with the concentrations of compound indicated. Compounds used were: G-744 (31) and P505-15 (32). For in vivo studies, G-744 and P505-15 were prepared as described in Supplemental Methods. Both compounds were dosed twice a day (BID) via oral gavage. Murine BR3-Fc or murine anti-ragweed control Abs (generated by Genetech, ref. 55), both containing a mIgG2a Fc, were dosed s.c. or i.p. at 5 mg/kg or 7.5 mg/kg 3 times per week as indicated. CTX (Baxter) was dosed at 30 mg/kg i.p. every 10 days.

B cell and monocyte/macrophage in vitro assays. Human B cells or monocytes were isolated from peripheral blood mononuclear cells (PBMCs) by Ficoll-Paque PLUS separation (GE Healthcare) and negatively selected by magnetic cell sorting as described (18). Murine splenic B cells were isolated by negative selection using the mouse B cell isolation kit (Miltenyi). Mouse BM-derived macrophages (BMM) were generated as described (18). Human and murine B cells were stimulated as described in Supplemental Methods. Methods for B cell pBtk, total Btk, pAkt and total Akt determination, CD86 expression, proliferation assays, Ca²⁺ flux, and survival are described in the Supplemental Methods. TNFα production by FcγR-activated purified monocytes or mouse BMM cultures was measured by ELISA (Bio-Rad) (18).

Plasmablast generation. Human memory B cells were differentiated into plasmablasts with in vitro stimulation conditions described in Supplemental Methods, followed by flow cytometry analysis at day 7.

Mice and in vivo BCR activation model. C57BL/6 or Balb/c mice were obtained from Charles River Laboratories; 8-week-old female NZB/W_F1 mice were from the Jackson Laboratory. Mice were housed under pathogen-free conditions until used in a study.

For BCR activation analysis, Balb/c mice were injected i.v. with 150 μg/mouse anti-IgD Abs (catalog GAM/IgD[Fc]/7S, Nordic-MUbios) 1 h after 1-time drug treatment. CD86 expression on splenic B220⁺ B cells was quantified 6 h after anti-IgD stimulation by flow cytometry.

AdIFNα accelerated NZB/W_F1 lupus model. LN was accelerated in 14- to 15-week old NZB/W_F1 mice by injecting i.v. 1.2 × 10⁸ IFNα rAd5-CMV (AdIFNα) pfu (Elizabeth Davis, Center for Cell & Gene Therapy, Baylor College of Medicine, Houston, Texas, USA) (55). This treatment typically results in elevated serum IFNα levels of 10 ng/ml 3 weeks after injection and 3 ng/ml 6 weeks after injection compared with LacZ rAd5-CMV control-treated mice. At 3 weeks after AdIFNα injection, mice were treated with vehicle or drugs.

Proteinuria and survival scoring. Proteinuria was determined weekly by colorimetric measurement using dipstick Multistix 10 SG on a Clinitek Status Analyzer (Siemens). Urine protein levels were scored as trace = 0, 30 mg/dl = 1, 100 mg/dl = 2, 300 mg/dl = 3, >300 mg/dl = 4, and death = 5. Progression-free survival was defined as the duration of remission, measured from the first time point when proteinuria was ≤ 300 mg/dl, or as survival time if there was no proteinuria progression.
Histology, histopathologic scoring, IHC, and immunofluorescence. To evaluate renal pathology and splenic GCs, kidneys and spleens were formalin fixed and paraffin embedded using routine methods. Renal IC deposition was determined by direct immunofluorescence staining on an autostainer (Dako Universal Autostainer) using Target antigen retrieval as per manufacturer's instructions followed by donkey anti–mouse IgG-AP48 (catalog A-21202) or goat anti–mouse IgM-AP48 (catalog A-21042, Invitrogen). Splenic GCs were detected by IHC on an autostainer (Dako) using Target antigen retrieval, followed by biotinylated anti–lectin-PNA (Vector Labs) and ABC-peroxidase Elite (Dako), and by chromogenic visualization with diaminobenzidine (Dako). All sections were visually evaluated in a blinded fashion by a board-certified pathologist using a Leica DM6000B microscope equipped with a Leica DFC500 camera and LAS software. Isotype or naive control Abs were used to confirm Ab specificity for all immunohistochemical and immunofluorescence stained tissues. Scoring systems for the different stains are described in Supplemental Methods.

Serum Ab ELISAs. Levels of total IgG, IgA, IgM, ANA, and anti-dsDNA Abs were determined by ELISA methodologies as described in detail in Supplemental Methods.

Flow cytometry. Single immune cell suspensions were prepared from spleen or peripheral blood after red blood cell lysis. Cells were counted on a FACS Calibur using Fluoresbrite Calibration Grade 6.0 micron YG microspheres (Polysciences), and flow cytometry staining was carried out as described in Supplemental Methods.

RNA sequencing and differential expression analyses. RNA from frozen spleen and kidney were isolated and analyzed by Illumina HiSeq and bioinformatics analysis performed as described in detail in Supplemental Methods. The raw RNA-sequencing reads and normalized data are available from GEO (accession number GSE72410).

Gene set enrichment analyses and identification of gene sets affected in spleen and kidney of LN mice. Immunology-related gene sets were derived from the immune response in silico (IRIS) microarray compendium and transcript modules from Chiche, et al. (39, 40). Significance for GSEA was determined as previously described (56). Gene set statistics were compared with gene set statistics derived from the same dataset with the sample labels permuted 499 times. Gene sets were considered significantly enriched if they showed a more extreme gene set statistic than > 99% of permutations (i.e. an empirical P value of 0.01).

Gene expression changes in spleen were analyzed using datasets published in ref. 41. To determine changes in gene expression that occur in specific kidney compartments and are regulated in proteinuric mice, we used 2 publicly available datasets: GSE32591 and GSE32583 (42). Bioinformatics methodology is described in Supplemental Methods. Lists of specific genes used for the various expression analyses in spleen and kidney are given in Supplemental Tables 2–6.

Statistics. Statistical analysis was performed using PRISM v6.0 (GraphPad). Comparison for each pair was performed using 2-tailed Student’s t test with Welch’s correction; multiple comparisons with a single control were performed using one-way ANOVA with Dunnett’s correction. Survival plots were analyzed by log-rank Mantel-Cox test. P < 0.05 were considered significant. Statistically significant differences are denoted in figures by glyphs. Absence of these notations indicates no significant differences. The IC50 for each assay was determined with PRISM by plotting the assay output versus the logarithm of the compound concentration using a nonlinear sigmoid with variable slope algorithm with the default curve-fitting parameters in the PRISM software.

Study approval. All animal studies were conducted in compliance with NIH Guide for the Care and Use of Laboratory Animals and were approved by the Genentech Institutional Animal Care and Use Committee.

Author contributions
AK, YW, and TH designed and performed experiments. JAH analyzed bioinformatics data. ES conducted lupus mice studies. YW and NR prepared RNA and performed RT-PCR analyses. CDA conducted the histology studies. AK and MB performed whole blood assays. JZC, LL, and HW performed PK analyses. KSC, PB, and WBY conceived the G-744 chemical series and synthesis. JJC facilitated G-744 production. JD and WPL supervised in vivo experiments. WPL established the lupus models. JAD and KR conceived the whole blood assays. JAD established the in vivo B cell CD86 assay. JH and JWL managed compound formulation. AJ supervised kinase and selectivity assays. JJC and AJ aided in manuscript assembly. JL performed in vivo B cell CD86 and survival studies. LED, SM, AN, CP, and JW performed Ab ELISA assays. ZL aided in FACS analyses. ML performed Akt phosphorylation assays. ZM generated RNA libraries and RNA sequencing data. MJT and KR conceived and analyzed the bioinformatics studies. KR designed and coordinated the research studies and analyzed the data. KR, MJT, and AK wrote the manuscript.
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