JAK Inhibitors in Inflammatory Bowel Disease: Recent Advances

Sailish Honap1,2, Alexandra Agorogianni1, Michael J Colwill1, Sonia Kalyanji Mehta1, Fiona Donovan1, Richard CG Pollock1,3, Andrew P Poullis1, Kamal V Patel1

1Department of Gastroenterology, St George's University Hospitals NHS Foundation Trust, London, UK

2School of Immunology and Microbial Sciences, King's College London, UK

3Institute of Infection and Immunity, St George's University, London, UK

Corresponding author and requests for reprints: Dr Sailish Honap, Department of Gastroenterology, 2nd Floor Grosvenor Wing, St. George’s Hospital, Blackshaw Road, London, SW17 0QT, UK. Email address: shonap@nhs.net, Telephone: 0208 725 1690

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# ABSTRACT (250 words)

Inflammatory bowel disease (IBD) commonly requires immunosuppressive treatments to induce and maintain durable remission. Janus kinase inhibitors (JAKI) are a novel group of orally administered, small molecule drugs that work by attenuating multiple cytokine signalling pathways to mediate dysregulated immune responses involved in the pathogenesis of IBD. Tofacitinib, filgotinib, and upadacitinib have demonstrated efficacy against placebo and are licensed for the treatment of moderate to severe ulcerative colitis; upadacitinib is the only JAKI also approved for the treatment of Crohn’s disease. Safety concerns stratified by age have led to class-wide regulatory restrictions for JAKI use across all inflammatory diseases. It is important for gastroenterologists managing patients with IBD to be aware of the key pivotal trial outcomes, to identify appropriate patients in whom to commence a JAKI, and to understand the safety considerations and ways to mitigate these risks in the patients they treat. This review provides a contemporaneous overview of this emerging therapeutic class and provides a practical guide for healthcare practitioners for initiating and monitoring JAKI in IBD.

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| **Key Points** |
| * Numerous pro-inflammatory cytokines involved in the pathogenesis of IBD signal through JAK-STAT pathways and are the therapeutic target for JAKI.
* JAKI offer a non-immunogenic, rapidly acting, oral option and should be considered for the treatment of moderate to severe IBD.
* Three JAKI are currently approved in the UK; tofacitinib and filgotinib are licensed for UC, while upadacitinib is licensed for both UC and CD.
* For safety reasons, JAKI should be used at the lowest dose possible and avoided in the following patients if no suitable alternative treatments are available; age ≥65 years, cigarette smokers or significant smoking history, risk factors for cancer, or risk factors for major adverse cardiovascular events (MACE).
* Further research is required to evaluate the efficacy of JAKI in acute severe UC, when used sequentially following prior JAKI failure, and when used in advanced combination therapy with a monoclonal antibody (mAb) to treat resistant disease/concomitant IMIDs.
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# JAKI: a novel class of orally administered drugs for the treatment of IBD

Inflammatory bowel disease (IBD), comprising ulcerative colitis (UC) and Crohn’s disease (CD), are immune-mediated inflammatory diseases (IMID) that typically require long-term immunosuppression. Broadly speaking, immunosuppressive therapies work in two main ways; to attenuate the signaling of one or more pro-inflammatory cytokines, or to prevent leukocyte migration to sites of inflammation. Treatment inefficacy and intolerance of existing therapies, however, has driven a proliferation of approved and investigational drugs over the past decade. JAK inhibitors (JAKI) are a novel class of drugs in IBD and were first introduced in 2019 with the approval of tofacitinib, followed more recently by the JAK1-selective filgotinib and upadacitinib (table 1). The drug development rationale for selective JAK inhibition is to maximise therapeutic efficacy without compromising the homeostatic functions of other JAK isoforms.

Janus kinases (JAK) are non-receptor tyrosine kinase proteins constitutively linked with the cytoplasmic domains of type I and type II cytokine receptors.1 The four JAK isoforms mediate cytokine signalling via the signal transducer and activator of transcription (STAT) pathway for more than 50 factors, including interleukins, interferons, hormones, and colony-stimulating factors that regulate a wide range of cellular processes.2,3 Inhibiting JAK phosphorylation and activation leads to a broad disturbance of cytokine signalling, including pathways pivotal to intestinal homeostasis and inflammation (figure 1). By corollary, this has safety implications and adverse events seen with tofacitinib and baricitinib for the treatment of rheumatoid arthritis have led to regulatory restrictions for all JAKI across the spectrum of immune-mediated inflammatory diseases.4,5

JAKI provide another treatment option for patients with moderate to severe IBD, particularly when failure and intolerance of existing therapies remains unacceptably high. JAKI are targeted, low molecular weight, synthetic drugs that hold a number of advantages over mAbs including, oral administration, quick absorption and a rapid onset of action, short half-life, and a lack of immunogenicity. This review provides a contemporary summary of the efficacy and safety data for the currently available JAKI to allow readers to take a balanced approach when considering this line of therapy for treating IBD patients.

# Tofacitinib: the first generation, non-selective JAKI licensed for UC

Tofacitinib (Xeljanz, Pfizer Inc.), is the first-in-class JAKI licensed for the treatment of moderate to severe UC. Originally developed as a JAK3 inhibitor to prevent solid organ transplant rejection, it is now known to be a pan-JAK inhibitor with preferential selectively for JAK1 and JAK3.6 Tofacitinib gained regulatory approval for UC in 2018 and is indicated for a total of five diseases including, rheumatoid arthritis (RA), psoriatic arthritis (PsA), juvenile idiopathic arthritis, and ankylosing spondylitis.7 Tofacitinib is not approved for CD after failing to meet the primary endpoint in the phase 2 trial.8 The OCTAVE UC phase III programme consisted of two identically designed eight-week induction studies (OCTAVE 1 and OCTAVE 2) and one maintenance 52-week study (OCTAVE SUSTAIN), which demonstrated superiority over placebo for clinical remission and several secondary endpoints.9 The more stringent definition of remission compared to earlier UC clinical trials had an additional requirement of a rectal bleeding sub score of 0. Table 2 summarises the registration trial data for approved JAKI in IBD.

Tofacitinib has been extensively studied since first entering the market for RA following Food and Drug Administration (FDA) approval in 2012.10 Prescription and sales data from the IQVIA databases estimate that more than half a million patients have been treated globally across all indications.11 For UC, long-term clinical trial and real world data demonstrate durable efficacy and highlight pertinent practice points, particularly with respect to dosing, that are not yet available for filgotinib and upadacitinib; safety outcomes are discussed later. In total, 1157 patients were studied through five randomised clinical trials, which includes OCTAVE Open, a long-term extension (LTE) study of 944 patients evaluating the safety and efficacy with up to seven years of treatment.12 Firstly, the LTE supports long term efficacy beyond one year. At three years, 59% and 34% of patients maintained or achieved clinical remission, with tofacitinib 5mg bd and 10 mg bd, respectively.12 This correlates with the relatively high degree of persistence seen with a median of 5.6 years for tofacitinib responders. Secondly, OCTAVE Open suggests healthcare practitioners should consider an extended induction for cases of sub-optimal response. Here, 52% of initial non-responders after eight weeks achieved a clinical response after an extended induction period totalling 16 weeks. At 36 months, 45% of delayed responders maintained clinical remission. Finally, most patients in stable remission on 10 mg bd maintenance therapy maintained remission following dose reduction to 5mg bd.13 Patients in deep endoscopic remission and those without prior anti-TNF-α failure were more likely to maintain remission on the lower dose. The wealth of real-world data that have been published are largely supportive of the efficacy data seen in the clinical trials. A recent meta-analysis of 17 studies and 1162 patients confirms tofacitinib effectiveness in a highly refractory patient population and showed pooled corticosteroid-free remission rates of 31% at one year.14

# Filgotinib: a JAK1-selective drug approved for UC but not CD

Filgotinib (Jyseleca, Galapagos NV) received MHRA licensing and NICE approval in 2022 for the treatment of moderate to severe UC after failure or intolerance of a biologic *or* conventional therapy.15,16 While filgotinib is also approved for the treatment of RA in the UK, European Union, and Japan, it is not available in the United States after a market authorisation application for RA was rejected by the FDA due to testicular toxicity concerns from pre-clinical studies. Gilead Sciences and Galapagos curtailed plans to pursue FDA authorisation despite subsequent safety studies showing no effect on semen parameters or sex hormones.17 Filgotinib is not approved for CD. Despite the favourable results of the phase II CD study (FITZROY), the induction cohorts of a large phase III study (DIVERSITY) of 1374 patients with moderate-to-severely active CD failed to meet the co-primary endpoints of endoscopic response and clinical remission at week 10.18,19 A phase II trial (DIVERGENCE 1) published in 2023 assessing the safety and efficacy of filgotinib in small bowel CD also did not show statistically significant differences against placebo.20

Filgotinib exhibits approximately 30-fold greater inhibition of JAK1 over JAK2- and other isoform-dependent signalling, which mechanistically is thought to maximise efficacy and minimise side effects.21,22 Efficacy in UC was demonstrated in the pivotal phase IIb/III SELECTION studies for UC, which were published in the *Lancet* in 2021.23 Biologic naïve and exposed patients were randomly assigned to receive once daily filgotinib at 100mg or 200mg or placebo in a 2:2:1 ratio across two 10-week induction studies; responders at the end of induction were then re-randomised into the 58-week maintenance study. The primary outcome was clinical remission as per the Mayo score defined by an endoscopic and stool frequency subscore of 0 or 1 and a rectal bleeding subscore of 0. The 200mg but not the 100mg od dose was efficacious at inducing and maintaining remission and met all pre-specified primary and secondary endpoints. Currently, there are no published real-world outcomes for the use of filgotinib in the treatment of UC but outcomes of the Galapagos-sponsored, prospective, observational study (GALOCEAN), are awaited.24

# Upadacitinib: a promising JAK1-selective drug and the first to be approved for both UC and CD

Upadacitinib (Rinvoq, AbbVie) is a second generation JAK1 selective inhibitor, and the only JAKI licensed for both moderate to severe UC and CD; MHRA licensing and NICE approval was received in 2023.25,26 Upadacitinib has six indications that also include RA, PsA, atopic dermatitis, axial spondyloarthritis.27 NICE recommends that upadacitinib is used for UC when conventional *or* biologic therapy has been ineffective or intolerable, similar to tofacitinib and filgotinib. For CD, however, patients must have had prior biologic exposure or have a contraindication to anti-TNF-α therapy prior to upadacitinib use.

Progression to the phase III trial programmes was based upon the success of the U-ACHIEVE and CELEST phase II dose-ranging studies for UC and CD, respectively.28,29 The phase III UC programme consisted of two replicate eight-week induction studies and a single 52-week maintenance study enrolling induction responders (table 2). The primary endpoint of clinical remission at both timepoints, based on an Adapted Mayo score that excluded Physician Global Assessment (PGA), was achieved with impressive superiority above placebo. Further, all secondary endpoints were met, which included clinical, endoscopic, histological, and quality of life (QoL) outcomes. Upadacitinib also improved faecal urgency and abdominal pain, symptoms that impact patient QoL but are rarely assessed in UC trials.

Marketing authorisation for upadacitinib in CD was supported by the results of the 12-week induction studies (U-EXCEL and U-EXCEED) including 1,021 patients, and a 52-week maintenance study (U-ENDURE) of 502 patients.30 Despite prior biologic exposure for three quarters of patients, drug efficacy assessed by the co-primary endpoints of clinical remission and endoscopic response at weeks 12 and 52 were striking. For induction, the proportion of patients in remission was nearly twice as high with upadacitinib compared to placebo and at least three times as high for endoscopic response. For maintenance, the delta above placebo was two to three-fold higher for remission versus placebo and almost six times higher for endoscopic response. Differences of this magnitude have not been seen in prior registration CD trials.

Results from the upadacitinib trials have generated optimism in the IBD community for several reasons. Firstly, and while between-trial efficacy comparisons are unwise, the marked difference seen above placebo across the phase III IBD programme compared with previous trials cannot be discounted, particularly given the stricter, endoscopy-incorporating endpoints. Secondly, in the absence of head-to-head studies, network meta-analyses have repeatedly ranked upadacitinib highest for the induction of remission in UC, and the highest for maintenance of remission in CD.31–33 Thirdly, the rapidity of onset, as significant improvement in symptoms were seen as early as 24 hours for UC and 14 days for CD.30,34 Finally, its efficacy in treating extraintestinal manifestations (EIM) and co-existing IMIDs. In U-ENDURE, upadacitinib was more likely than placebo to resolve EIMs at the 30mg dose and among all JAKI, upadacitinib has the highest number of therapeutic indications for IMIDs. The U-ENDURE LTE and the prospective real-world studies PROFUNDUS (UC) and UPlift (CD) will determine the durability of efficacy and safety and its generalisation to everyday practice.35,36

# Table 1: Currently licensed JAK inhibitors for inflammatory bowel disease

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| --- | --- | --- | --- |
|  | Tofacitinib (Xeljanz) | Filgotinib (Jyseleca) | Upadacitinib (Rinvoq) |
| Manufacturer | Pfizer | Gálapagos | AbbVie |
| Selectivity | JAK3 >JAK2 >JAK1 | JAK1 >JAK2 >JAK3 and TYK2 | JAK1 >JAK2 and JAK3 |
| Ulcerative colitis | MHRA/FDA/EMA licensedNICE approved | MHRA/EMA licensedNICE approved | MHRA/FDA/EMA licensedNICE approved |
| Crohn’s disease | Unlicensed | Unlicensed | MHRA licensedNICE approved |
| Induction dose | 10mg bd for 8-16 weeks | 200mg od for 10-22 weeks | 45mg od for 8-16 weeks for UC45mg od for 12 weeks for CD |
| Maintenance dose | 5mg bd | 200mg od | 15-30 mg od |
| Drug half life | 3 hours | 7 hours | 9-14 hours |
| Metabolism | 65% Hepatic [CYP3A4 and CYP2C19] | Intestinal [CES2 [primarily]] and Hepatic [CES1] | 34% Hepatic [CYP3A4 and CYP2D6] |
| Liver disease | Avoid Child-Pugh C | Avoid Child-Pugh C | Avoid Child-Pugh C |
| Renal disease | ↓ dose if CC <30mL/min | ↓ dose if CC <60mL/min | ↓ dose if CC <30mL/min |
| Concomitant IMM studied | Not studied for IBD | Thiopurine and methotrexate | Methotrexate |
| Additional Indications | RA, PsA, AS, JIA | RA | RA, PsA, AS, AD |

Abbreviations: JAK, Janus kinase; MHRA, Medicines and Healthcare products Regulatory Agency; FDA, Food and Drug Administration; EMA, European Medicines Agency; NICE, National Institute for Health and Care Excellence; bd, twice daily; od, once daily; CYP, cytochrome P450; CES, Carboxylesterase; CC, creatinine clearance; IMM, immunomodulator; RA, rheumatoid arthritis; PsA, psoriatic arthritis; AS, ankylosing spondylitis; JIA, juvenile idiopathic arthritis; AD, atopic dermatitis

# Table 2: Summary of pivotal trials for licensed JAK inhibitors in inflammatory bowel disease

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| --- | --- | --- | --- | --- | --- | --- | --- |
|  |  | Trials | Treatment Phase | n | Duration (w) | Primary Objective | Key Findings |
| **UC** | **Tofacitinib**9 | OCTAVE 1 | Induction | 598 | 8 | Clinical remission\* at week 8 | Clinical remission in 18.5% (tofacitinib 10mg bd) vs. 8.2% (placebo), p=0.007. |
| OCTAVE 2 | Induction | 541 | 8 | Clinical remission at week 8 | Clinical remission in 16.6% (tofacitinib 10mg bd) vs. 3.6% (placebo), p<0.001. |
| OCTAVE SUSTAIN | Maintenance | 593 | 52 | Clinical remission at week 52 | Clinical remission in 34.3% (tofacitinib 5mg bd) vs. 40.6% (tofacitinib 10mg bd), vs. 11.1% (placebo), p<0.001 for both comparisons with placebo. |
| **Filgotinib**23 | SELECTIONSTUDY A | Induction | 659 | 10 | Clinical remission at week 10 | Clinical remission in 26.1% (filgotinib 200mg od) vs. 15.3% (placebo), p=0·0157. |
| SELECTIONSTUDY B | Induction | 689 | 10 | Clinical remission at week 10 | Clinical remission in 11.5% (filgotinib 200mg od) vs. 4.2% (placebo), p=0·0103. |
| SELECTION | Maintenance | 664 | 58 | Clinical remission at week 58 | Clinical remission in 37.2% (filgotinib 200mg od) vs. 11.2% (placebo), p<0·0001. Clinical remission was not significantly different between filgotinib 100mg od and placebo at week 10 but was significant by week 58 (23·8% vs.13·5%, p=0·0420). |
| **Upadacitinib**37 | U-ACHIEVE | Induction | 474 | 8 | Clinical remission at week 8 | Clinical remission in 26% (upadacitinib 45mg od) vs 5% (placebo), p<0.0001. |
| U-ACCOMPLISH | Induction | 522 | 8 | Clinical remission at week 8 | Clinical remission in 34% (upadacitinib 45mg od) vs 4% (placebo), p<0.0001. |
| U-ACHIEVE | Maintenance | 451 | 52 | Clinical remission at week 52 | Clinical remission in 52% (upadacitinib 30mg od), 42% (upadacitinib 15mg od), 12% (placebo), p<0.0001. |
| **CD** | **Upadacitinib**30 | U-EXCEL | Induction | 526 | 12 | Clinical remission and endoscopic response at week 12\*\* | Clinical remission in 49.5% (upadacitinib 45mg od) vs. 29.1% (placebo) and endoscopic response in 45.5% (upadacitinib 45mg od) vs. 13.1% (placebo). P<0.001 for both comparisons. |
| U-EXCEED | Induction | 495 | 12 | Clinical remission and endoscopic response at week 12 | Clinical remission in 38.9% (upadacitinib 45mg od) vs. 21.1% (placebo) and endoscopic response in 34.6% (upadacitinib 45mg od) vs. 3.5% (placebo). P<0.001 for both comparisons. |
| U-ENDURE | Maintenance | 502 | 52 | Clinical remission and endoscopic response at week 52 | Clinical remission in 47.6% (upadacitinib 30mg od) vs. 37.3% (upadacitinib 15mg od) vs 15.1% (placebo) and endoscopic response in 40.1% (upadacitinib 30mg od) vs. 27.6% (upadacitinib 15mg od) vs. 7.3% (placebo). P<0.001 for all comparisons. |

Abbreviations: CD, Crohn’s disease; n, number of patients; UC, ulcerative colitis; w, weeks. \*Definition of clinical remission was similar across the UC trials and was defined as a total Mayo score of ≤2, with no subscore >1 and a rectal bleeding subscore of 0. For the filgotinib studies, a 1-point decrease in stool frequency from induction baseline for a subscore of 0 or 1 was required and for the upadacitinib studies, the Physician Global Assessment was removed due to subjectivity. \*\*Remission defined as CD activity index <150, and endoscopic response defined as a fall in Simple Endoscopic Score for CD score >50% from baseline.

# SAFETY OF JAK INHIBITORS

## General safety considerations

An important barrier to initiating any immunosuppressive therapy is the concern of adverse events from patients and clinicians. Most adverse events related to JAKI are mild to moderate, predictable, and easy to manage. However, JAKI are also associated with more serious side effects. The randomised, open-label, noninferiority, safety study of tofacitinib (ORAL SURVEILLANCE) in a cardiovascular risk-enriched population showed increased incident rates (IR) of cancer and MACE when compared to anti-TNF-α therapy; similar FDA-mandated studies of baricitinib in RA are ongoing and are estimated to complete enrolment by 2025.38–40 It is difficult to ascertain with any certitude whether certain adverse events are class-specific, drug-specific, or patient/population-specific. Figure 2 outlines how to mitigate and manage some of the potential adverse events from JAKI and figure 3 highlights important pre- and post-initiation considerations of JAKI treatment and describes situations where JAKI are best avoided.41

At baseline, all patients should have a pre-immunosuppression screen for infection and be up to date with recommended vaccinations in accordance with BSG and ECCO guidelines.42,43 JAKI approved for IBD should be avoided in patients with advanced liver disease (Child-Pugh C), particularly tofacitinib and upadacitinib that are predominantly metabolised by hepatic CYP3A4. Filgotinib is metabolised by carboxylsterase-2 (CES2), is renally excreted, and the creatinine clearance threshold for dose reduction is lower than other JAKI (table 1). Laboratory parameters should be monitored after induction and throughout maintenance therapy as these have been shown to be affected by JAKI. Abnormalities in blood counts and liver transaminases in the short and long-term were mild to moderate and included anaemia, leukopenia, and elevations in aminotransferases. Most patients experiencing these abnormalities were able to remain on the study drug and for patients where JAKI discontinuation was necessary, these changes resolved.12,44,45

## Major adverse cardiovascular events

Chronic systemic inflammation is known to promote accelerated atherosclerosis and is a risk factor for MACE, defined by a fatal or non-fatal myocardial infarction or ischaemic stroke. In IBD patients, this risk is modestly increased, particularly during active disease.46–48 The JAKI class are also associated with dose-dependent increases of HDL and LDL cholesterol without affecting the HDL:LDL ratio, changes that are reversible on treatment cessation.49 When compared to anti-TNF-α agents, tofacitinib has been shown to increase the incidence of MACE in RA patients aged ≥50 years with at least one cardiovascular risk factor, hazard ratio (HR) 1.33 (95% confidence interval (CI), 0.91 to 1.94).38 It is important to note that the increased risk of MACE with JAKI has not been demonstrated in patients with IBD and the overall incidence is low.50 Nevertheless, steps should be taken to modify cardiovascular risk factors such as lipid-lowering and smoking cessation, and if no other treatments exist, JAKI should be avoided in high-risk populations.

## Venous Thromboembolism

IBD patients with active disease have a twofold increased risk of venous thromboembolism (VTE).51 Cases of VTE have been observed with tofacitinib, filgotinib, and upadacitinib in the trial programmes and although not powered to assess VTE, there was an increased number of VTE events in ORAL SURVEILLANCE, particularly with the 10 mg bd tofacitinib dose. However, two large meta-analyses of 5,143 and 6,542 JAKi-exposed patients across all IMIDs, did not find an increased risk.52,53 It remains unclear whether JAKI have a direct causal role in the development of VTEs or whether these VTEs occur in the context of a higher baseline risk in IMIDs. For now, the regulatory guidance is to use JAKI with caution in patients with VTE risk factors.4,5

## Cancer including non-melanoma skin cancer (NMSC)

Uncontrolled inflammation is a critical component of the neoplastic process, driving cell proliferation, survival, and migration.54 Compared to the general population, IBD may be associated with an increased risk of overall cancer and cancer-specific mortality.55

ORAL SURVEILLANCE did not meet its pre-specified non-inferiority criteria and tofacitinib was associated with an increased cancer incidence (excluding NMSC) compared to anti-TNF-α agents, HR 1.48 (95% CI, 1.04 to 2.09).38 In a recent meta-analysis of incorporating 62 RCTs, 16 LTE studies, and 82,366 person-years of JAKI exposure, risk of malignancy did not differ significantly between JAKI and placebo but was associated with a higher incidence of malignancy when compared to anti-TNF-α agents, largely influenced by ORAL SURVEILLANCE.56 JAKI-induced cancers remain a rare occurrence but until this risk is precisely elucidated, JAKI should be avoided in patients with an increased cancer risk, such as those with an active or past cancer.

Aggressive squamous cell carcinomas have been reported with JAKI; with previous NMSC, prior anti-TNF failure, and older age, associated with increased NMSC risk.57,58 Manufacturer guidance for JAKI suggest periodic skin examination, particularly those at higher risk of skin cancer. Reassuringly, however, the IR of NMSC across the clinical trial programmes for JAKi were low at 0.51/100 patient-years; IR among patients exposed to comparator (mainly placebo) was 0.27/100 patient-years.

## Infection

The commonest infections affecting the IBD-approved JAKI in the trials were nasopharyngitis, upper respiratory tract infections, and influenza.9,23,37,30 The IR of serious infections across all JAKI was 2.81/100 person-years and were mainly bacterial, including pneumonias, urinary tract and skin infections.53 Rates of tuberculosis were low in regions of low to medium incidence but should be screened for pre-initiation.59 The most notable viral infection was a dose-dependent increased risk of herpes zoster infection, which has been associated across the JAKI class. This may be explained by the JAK1-mediated suppression of interferon, which has an antiviral role.60 Most cases are non-serious affecting a single dermatome and did not lead to treatment withdrawal. Using the lowest JAKI dose possible and vaccination against varicella zoster reactivation are the best ways of mitigating this risk. Live vaccines are contraindicated during JAKI therapy but the inactivated shingles vaccine (Shingrix) is to be made available in the UK later in 2023 for immunocompromised patients aged ≥50 where previously it was reserved for those aged ≥70 years.61

## Pregnancy and breastfeeding

JAKI are small molecule drugs that cross the placenta throughout pregnancy and have recently been shown to be excreted into human breast milk.62 Pre-clinical studies have shown that JAKI at doses much higher than those approved for humans were teratogenic and feticidal.7,16,63 Contraception is advised for all female patients during treatment JAKI and for one week after the last dose of filgotinib and four weeks for tofacitinib and upadacitinib, as per manufacturer instructions. Breastfeeding is not recommended. Outcomes from maternal tofacitinib exposure of 62 patients across the UC, RA, and psoriasis trials were not dissimilar to the general population, although this is hindered by a small sample size.64,65 Pregnancy outcomes for filgotinib and upadacitinib are scarce.

## Restrictions to JAKI use: update from the EMA and MHRA

Following the publication of ORAL SURVEILLANCE in early 2022, the JAKI class of drugs were placed under safety review by the Pharmacovigilance Risk Assessment Committee (PRAC), a branch of the EMA.4 PRAC recommendations were issued in October 2022 and were endorsed and approved by the EMA’s European Commission in March 2023.4 The MHRA followed suit and issued a statement in April 2023.5 Box 1 outlines these recommendations, which apply to all JAKI indicated for chronic inflammatory diseases; fedracitinib (Inrebic) and ruxolitinib (Jakavi) for myeloproliferative diseases, and baricitinib, when used short-term for the treatment of COVID-19, are exempt. There is a degree of clinical judgement with the wording of the recommendations, for example, the degree of previous smoking exposure and risk factors for cancer are open to interpretation. This is unlikely to be the end of the turbulent JAKI story; while filgotinib and upadacitinib are exempt from post-market FDA-required studies, results from the ongoing baricitinib studies may well lead to yet another change in the recommendations and guidance.

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| **Box 1: EMA and MHRA Recommendations on JAK Inhibitors Use** |
| JAKI only to be used in the following groups at the lowest effective dose if no suitable agent available:* aged ≥65 years
* at increased risk of major cardiovascular problems (such as heart attack or stroke)
* smokers or have smoked for a long time in the past
* increased risk of cancer

JAKIs to be used with caution in those with VTE risk factors and at the lowest effective dose |

# FUTURE PERSPECTIVES AND CONCLUSIONS

There are several pan-JAK and JAK-selective inhibitors, including TYK2 inhibitors, in various stages of development for IBD.66–68 Even with the three available options, how do we select between therapies in this increasingly crowded space? While many unanswered questions remain, factors that will guide decision making will likely centre on reimbursement cost, available direct and indirect safety and efficacy data, number of licensed indications to treat EIMs/co-existing IMIDs, clinician familiarity, and dosing regimens; filgotinib and upadacitinib are dosed once daily as the prolonged release tofacitinib tablets are unavailable in the UK. The limelight on tofacitinib’s safety profile has led to the perception that JAK-1 selective inhibitors are safer. Whether selective inhibition of a JAK isoform confers a better safety and efficacy profile remains to be seen; filgotinib and upadacitinib will not be subjected to the same FDA-mandated testing as tofacitinib and baricitinib were so longer term studies are essential.69 Theoretically, selectivity of JAK isoform inhibition could limit side effects, although this selectivity is dose and tissue dependent, and may be lost with increasing doses.70 Key areas of further research with direct clinical relevance include, JAKI use in advanced combination therapy together with a mAb, JAKI positioning and sequencing after prior JAKI failure/intolerance, mechanisms of resistance and loss of response, and its role in the treatment of acute severe UC.

JAKI are an emerging group of highly efficacious drugs used to treat inflammatory, autoimmune, and myeloproliferative diseases. For IBD, they represent another key line of therapy for patients with active disease and their advantages over mAbs, the crux of IBD pharmacological therapy over the past two decades, has attracted widespread attention. The safety profile of this drug class and the regulatory restrictions in place for all JAKI for inflammatory diseases require careful benefit-risk consideration when initiating and monitoring therapy. Although these risks in absolute terms are likely to be small for the typical IBD patient population, they are not inconsequential. Personalised stratification to appropriate JAKi use is an attractive future goal.

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# TABLE LEGENDS

## Table 1: Overview of licensed JAK inhibitors for IBD

## Table 2: Summary of pivotal phase III trials for JAKI in IBD

# FIGURE LEGENDS

## Figure 1: The structure of a JAK protein and the mechanism of action of JAK inhibitors

a) *Functional domains of a JAK protein.* This schematic displays the four functional domains common to all JAKs. The kinase domain is responsible for enzymatic activity and substrate phosphorylation while the pseudokinase domain lacks enzymatic activity but regulates the active catalytic kinase domain and is also associated with STAT recruitment. The Src-homology-2 (SH2) and FERM domains are involved in associating the JAK with the respective cell membrane receptors. Alternative nomenclature in the literature describes seven JAK homology domains (JH) numbered from the carboxyl terminus to the amino terminus.

b) *JAK-STAT signal transduction and the mechanism of action of JAKI in attenuating cytokine signalling.* The four members of the JAK family, JAK1, JAK2, JAK3, and Tyk2, are located on the intracellular domain of the cell surface receptor and are essential in mediating signalling downstream of cytokine receptors for a broad range of cellular processes. Numerous cytokines implicated in the pathogenesis of IBD, involved in both innate and adaptive immunity, signal through this pathway. Different cytokine-activated JAK-STAT combinations drive different cellular processes with a high degree of specificity. Extracellular cytokine binding result in conformational receptor changes, which brings the associated JAKs in close proximity and in turn, results in auto- and transphosphorylation of receptor chains. These forms docking sites for downstream STAT proteins, which after phosphorylation, receptor chains dissociation, and STAT homo- or hetero- dimerisation, translocate to the nucleus to modulate gene transcription. JAKI reversibly bind onto the adenosine triphosphate (ATP) site on the catalytic cleft of the kinase domain to prevent JAK phosphorylation and activation and inhibits the ensuing signalling cascade.

## Figure 2: Managing potential adverse events of JAK inhibition

Abbreviations: ALC, absolute lymphocyte count; ANC, absolute neutrophil count; CVS, cardiovascular; CVD, cerebrovascular disease; ESRF, end stage renal failure; MACE, major adverse cardiovascular events; IHD, ischaemic heart disease; LFTs, liver function tests; VTE, venous thromboembolism.

## Figure 3: A checklist for initiating and monitoring therapy

Abbreviations: BSG, British Society of Gastroenterology; FBC, full blood count; JAKI, Janus kinase inhibitor; LFT, liver function tests; MACE, major adverse cardiovascular events; VTE, venous thromboembolism; \*if no suitable alternative available.