Safety profile of upadacitinib over 15 000 patient-years across rheumatoid arthritis, psoriatic arthritis, ankylosing spondylitis and atopic dermatitis

Gerd R Burmester,1 Stanley B Cohen,2 Kevin L Winthrop,3 Peter Nash,4 Alan D Irvine,5,6 Atul Deodhar,3 Eduardo Mysler,7 Yoshiya Tanaka,8 John Liu,9 Ana P Lacerda,9 Hannah Palac,9 Tim Shaw,10 Philip J Mease11, Emma Guttman-Yassky12

ABSTRACT

Objective To evaluate the long-term safety profile of upadacitinib across rheumatoid arthritis (RA), psoriatic arthritis (PsA), ankylosing spondylitis (AS) and atopic dermatitis (AD).

Methods Safety data from clinical trials of upadacitinib 15 mg and upadacitinib 30 mg (AD only) for treating RA, PsA, AS and AD as of 30 June 2021 were analysed; some RA and PsA studies included adalimumab and methotrexate as active comparators. Treatment-emergent adverse events (TEAEs) were presented by disease as exposure-adjusted event rates per 100 patient years (E/100 PY).

Results The analysis included 6991 patients (RA, n=3209; PsA, n=907; AS, n=182; AD, n=2693) who received at least one dose of upadacitinib, representing 15 425 PY of exposure (maximum duration 2.75–5.45 years) across diseases. Rates (E/100 PY) of any TEAE (205.5–278.1) and TEAE leading to discontinuation (4.5–5.4) were similar across diseases; serious TEAEs were numerically higher in patients with RA and PsA. Rates of herpes zoster (1.6–3.6), non-melanoma skin cancer (0.3–1.4) were observed, with rates generally lowest in AD. Rates of infections, elevations in creatine phosphokinase levels (4.4–7.9) were higher with upadacitinib than with active comparators in the RA and PsA populations. Deaths (0–0.8), serious infections (0–3.9), major adverse cardiovascular events (0–0.4), venous thromboembolism (<0.1–0.4) and malignancies (0.3–1.4) were observed, with rates generally lowest in AD and AS. Increased rates of acne were observed in patients with AD only.

Conclusions Findings from this analysis demonstrate that upadacitinib is generally well tolerated with observed differences in safety profiles likely reflective of varying patient characteristics across RA, PsA, AS and AD populations.

Trial registration numbers NCT02675426, NCT02706951, NCT02706847, NCT02629159, NCT02706873, NCT03086343, NCT03104374, NCT03104400, NCT03178487, NCT03569293, NCT03568318 and NCT03607422.

INTRODUCTION

Immune-mediated inflammatory diseases (IMIDs), including rheumatoid arthritis (RA) and spondyloarthritides, are common,
clinically diverse and chronic. While significant variations in population distribution, tissue localisation, clinical phenotypes and therapeutic responses are apparent between different IMIDs, they share many similar pathophysiological mechanisms. Small-molecule Janus kinase (JAK) inhibitors are a therapeutic class for the management of IMIDs. Although certain adverse events (AEs) appear associated with approved JAK inhibitors, including higher rates of herpes zoster and elevation in creatine phosphokinase (CPK) levels, the safety of individual JAK inhibitors continues to be evaluated in diverse patient populations impacted by each of these chronic conditions.

Recent outcomes of the Oral Rheumatoid Arthritis trial—L (ORAL) Surveillance study, comparing the JAK inhibitor tofacitinib to tumour necrosis factor (TNF) inhibitor therapy, further highlight the need to characterise the safety profile of JAK inhibitors, especially in the context of active comparators.

Upadacitinib is an oral, reversible JAK inhibitor that has been and continues to be studied in comprehensive phase III clinical programmes including trials in RA, psoriatic arthritis (PsA), ankylosing spondylitis (AS), atopic dermatitis (AD) and ulcerative colitis (UC). While 15mg once a day (QD) is the approved dose for treating rheumatological diseases, upadacitinib 15 and 30mg (if response is inadequate with 15mg) QD are approved for treating moderate-to-severe AD in adults and adolescents, and upadacitinib 45mg was recently approved in the USA for treating patients with UC as induction therapy for 8 weeks followed by QD upadacinib 15mg or 30mg as maintenance therapy. Here, we report an integrated analysis of the safety profile of QD upadacitinib across indications approved as of December 2021 including RA, PsA, AS and AD, along with active comparator data from trials including adalimumab or methotrexate (MTX).

METHODS
Patients and studies
This analysis included safety data from 12 upadacitinib clinical trials (online supplemental table S1) with a cutoff date of 30 June 2021 for all studies. Studies of RA, PsA and AS included patients aged ≥18 years, while the studies of AD included adult and adolescent (aged ≥12 years) patients.

Patient involvement
Patients and/or the public were not involved in the design, conduct, reporting or dissemination plans of this research.

Dosing
Data are pooled across studies within each disease and reported for the following treatment groups: upadacitinib 15mg QD (RA, PsA, AS and AD), upadacitinib 30mg QD (AD only, the only included indication approved at the 30mg dose), adalimumab 40mg subcutaneously every other week (RA [SELECT-COMPARE study], PsA [SELECT-PsA 1 study] and MTX [RA, SELECT-EARLY study]). Depending on study protocol, patients receiving upadacitinib did so either alone as monotherapy or in combination with MTX or other conventional synthetic disease-modifying antirheumatic drug (csDMARD) therapy. In SELECT-COMPARE, all patients, including those receiving upadacitinib and adalimumab, received background MTX; in SELECT-PsA 1, patients could have received adalimumab with or without concomitant background csDMARDs. All MTX data come from SELECT-EARLY (RA) and reflect monotherapy use.

Safety assessments
We report an overview of treatment-emergent adverse events (TEAEs) and adverse events of special interest (AESIs) to the JAK inhibitor class. TEAEs are defined as onset on or after the first dose of study drug and up to 30 days after the last dose of upadacitinib or MTX or up to 70 days after the last dose for adalimumab. Non-TEAEs are included only for deaths. An additional subanalysis is provided for TEAEs in patients receiving upadacitinib as monotherapy or in combination with any csDMARD at baseline for RA and PsA only.

TEAEs were coded using the Medical Dictionary for Regulatory Activities System Organ Classes and Preferred Terms. All deaths, potential cardiovascular (CV) events and arterial and venous thromboembolisms (VTEs) were adjudicated by a blinded, independent cardiovascular adjudication committee. Gastrointestinal perforations were blindly adjudicated by sponsor-employed experts independent of the upadacitinib clinical programmes using a prespecified definition of acute gastrointestinal tract perforation, which is non-iatrogenic and non-traumatic.

Major adverse cardiovascular events (MACEs) included CV death, non-fatal myocardial infarction and non-fatal stroke. VTE included deep vein thrombosis (DVT) and pulmonary embolism (PE). CV risk factors included history of a CV event, hypertension or diabetes mellitus; current or former tobacco use; elevated low-density lipoprotein cholesterol (>3.36mmol/L) levels or lowered high-density lipoprotein cholesterol (<1.034mmol/L) levels. Active tuberculosis and herpes zoster are separately assessed from other opportunistic infections. Laboratory-related abnormalities (anaemia, neutropenia, lymphopenia, hepatic disorder and elevated CPK levels) are based on investigator-reported AEs. Potentially clinically significant values (grade 2, 3 and 4 changes) for select laboratory parameters of interest to the JAK inhibitors class are also provided. For RA, the toxicity grading scales are based on OMERACT (Outcome Measures in Rheumatoid Arthritis Clinical Trials) criteria, with the exception of creatine kinase and creatinine, which are based on NCI Common Terminology Criteria for Adverse Events (CTCAE). For PsA, AS and AD, all toxicity grading scales are based on CTCAE.
Statistical analyses

All TEAEs are reported as exposure-adjusted event rates (EAEs, exposure-adjusted event rates per 100 patient years [E/100 PY]) based on the treatment received at the time of the AE with 95% CIs calculated using the exact method for the Poisson mean; all events, including multiple events occurring in a single patient, were included in the numerator. For EAEs, exposure time was calculated as the total study drug duration. We also calculated exposure-adjusted incidence rates (EAIrS) with 95% CIs using the exact method for the Poisson mean for AESIs (online supplemental material); EAIrS were calculated as the number of patients with one or more event per 100 PY (n/100 PY). In patients who experienced an event, exposure time was calculated as the time to the first event; in patients who did not experience an event, exposure time was censored on the day of the patient’s last assessment or cut-off date of database lock, whichever occurred first. Additional subanalyses include EAER stratified by 6-month intervals (reported for MACE, VTE, malignancy excluding non-melanoma skin cancer (NMSC), serious infections and herpes zoster) and EAEs for all AESIs stratified by age group (patients aged <65 years and ≥65 years).

Standardised mortality ratio (SMR) was determined using the WHO country-specific, age-specific and gender-specific death data for the general population; 95% CIs were calculated using Byars’ approximation, and calculations included COVID-19 deaths. Standard incidence ratio (SIR) for malignancy excluding NMSC was determined using age-gender-adjusted data from the Surveillance, Epidemiology, and End Results (SEER) cancer incidence rate data in 2000–2018 from the US general population and adjusted for age at baseline; 95% CIs were calculated following a Poisson distribution.

RESULTS

Patients and exposure

Patient data collected from 12 studies included 3209 patients with RA (PY=9079.1), 907 with PsA (PY=1872.3), 182 with AS (PY=320.1) and 2693 with AD (PY=2035.8 for upadacitinib 15 mg and 2118.0 for upadacitinib 30 mg). The maximum duration of treatment was longest in the RA programme (5.45 years, median 3.46 years) and shortest in the AD programme (2.75 years, median 1.62 years). Reference comparator arms included patients receiving at least one dose of adalimumab (RA: 579 patients, 1307.7 PY; PsA: 429 patients, 903.7 PY) and patients receiving at least one dose of MTX (314 patients with RA only, 781.7 PY). The majority of patients receiving upadacitinib also received concomitant csDMARD therapy in RA (2548 patients) and PsA (642 patients), whereas concomitant csDMARD use was infrequent in patients with AS and AD (table 1). Other noteworthy differences in baseline characteristics typical of the respective disease state were present, including younger patients in the AD population and a higher proportion of female patients in the RA population (table 1). Across all groups, 48.5%–83.5% of patients had at least one CV risk factor at baseline.

Overview of AEs

Rates of any TEAEs with upadacitinib ranged from 205.5 with upadacitinib 15 mg in RA to 278.1 with upadacitinib 30 mg in AD (table 2). In RA, the rates were generally comparable between upadacitinib and active comparators and in patients receiving upadacitinib as monotherapy or in combination with csDMARD therapy (online supplemental table S2); in PsA, the rate was numerically higher with upadacitinib (244.8) vs adalimumab (229.9). The highest rates of serious TEAEs were observed in RA. TEAEs leading to discontinuation were generally comparable across all treatment groups and diseases. In AD, rates of any TEAE, serious TEAE and TEAE leading to discontinuation were numerically higher with upadacitinib 30 mg compared with upadacitinib 15 mg. The incidence of deaths was <1.0/100 PY in the RA programme and similar across upadacitinib, adalimumab and MTX groups. In PsA, rates of death were higher with upadacitinib 15 mg compared with adalimumab, owing to increased COVID-19-related deaths in patients taking upadacitinib. No deaths were reported in AS, and three deaths were reported in the upadacitinib 30 mg AD group, with two of the three deaths related to COVID-19. Excluding COVID-19, the most common cause of death was related to CV disease, followed by other infections and malignancies. The SMR estimates for each disease were 0.59 (95% CI: 0.43 to 0.78) for RA, 0.59 (95% CI: 0.36 to 0.92) for PsA and 0.28 (95% CI: 0.06, 0.82) for AD (both doses), with no evidence suggesting that the number of deaths in patients with RA, PsA or AD exposed to upadacitinib were higher than what would have been expected for the general population.

In RA, PsA, AS and AD populations, the most common TEAE associated with upadacitinib 15 mg was upper respiratory tract-related infection (online supplemental table S3), while acne was the most common TEAE associated with upadacitinib 30 mg in AD. Acne was infrequently reported in the rheumatological diseases; while events of acne in AD were generally mild or moderate, non-serious and rarely led to treatment discontinuation. The pattern, characteristics and incidence of COVID-19 infections, including frequency of events and those leading to hospitalisation, observed in patients receiving upadacitinib were generally similar to what has been observed in the general population (online supplemental table S4).

Adverse events of special interest

AESIs are presented in figure 1 in EAERs and in the online supplemental figure S1 in EAIRs.

Serious and opportunistic infections

Serious infections with upadacitinib occurred at similar rates between RA, PsA and AD and infrequently led to discontinuation; no serious infections were reported in
<table>
<thead>
<tr>
<th>Parameter, n (%)</th>
<th>RA</th>
<th>PsA</th>
<th>AS</th>
<th>AD</th>
</tr>
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<tbody>
<tr>
<td><strong>Parameter, n (%)</strong></td>
<td><strong>UPA 15mg QD N=3209</strong></td>
<td><strong>ADA 40mg EOW N=579</strong></td>
<td><strong>MTX N=314</strong></td>
<td><strong>UPA 15mg QD N=907</strong></td>
</tr>
<tr>
<td>Age (years), mean (SD)</td>
<td>54.3 (12.0)</td>
<td>54.1 (11.7)</td>
<td>53.2 (12.9)</td>
<td>51.5 (12.1)</td>
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<td>Age ≥65 years</td>
<td>643 (20.0)</td>
<td>106 (18.3)</td>
<td>58 (18.5)</td>
<td>129 (14.2)</td>
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<td>Female sex</td>
<td>2581 (80.4)</td>
<td>470 (81.2)</td>
<td>240 (76.4)</td>
<td>518 (57.1)</td>
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<tr>
<td>BMI, mean (SD)</td>
<td>29.1 (6.7)</td>
<td>29.4 (7.1)</td>
<td>28.0 (6.3)</td>
<td>30.6 (6.9)</td>
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<td>BMI ≥30 kg/m²</td>
<td>1200 (37.4)</td>
<td>227 (39.2)</td>
<td>97 (30.9)</td>
<td>429 (47.3)</td>
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<td>Time since diagnosis (years), mean (SD)</td>
<td>8.5 (8.4)</td>
<td>8.2 (8.0)</td>
<td>2.6 (5.1)</td>
<td>7.2 (7.8)</td>
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<td>Disease activity</td>
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<td></td>
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<td>CRP-based disease measure, mean (SD)*</td>
<td>5.8 (1.0)</td>
<td>5.9 (1.0)</td>
<td>5.9 (1.0)</td>
<td>57.5 (29.7)</td>
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<td>EASI, mean (SD)</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
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<td>Concomitant therapies</td>
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<td>csDMARD(s)</td>
<td>2548 (79.4)</td>
<td>579 (100)</td>
<td>0</td>
<td>642 (70.8)</td>
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<td>MTX alone</td>
<td>2180 (67.9)</td>
<td>579 (100)</td>
<td>0</td>
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<td>csDMARD other than MTX</td>
<td>198 (6.2)</td>
<td>0</td>
<td>0</td>
<td>89 (9.8)</td>
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<td>Corticosteroids</td>
<td>1761 (54.9)</td>
<td>350 (60.4)</td>
<td>164 (52.2)</td>
<td>134 (14.8)</td>
</tr>
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<td>Topical glucocorticoids</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
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<td>NSAIDs</td>
<td>2034 (63.4)</td>
<td>362 (62.5)</td>
<td>223 (71.0)</td>
<td>566 (62.4)</td>
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<td>Aspirin</td>
<td>269 (8.4)</td>
<td>36 (6.2)</td>
<td>24 (7.6)</td>
<td>89 (9.8)</td>
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<td>Statin use</td>
<td>369 (11.5)</td>
<td>55 (9.5)</td>
<td>26 (8.3)</td>
<td>123 (13.6)</td>
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<td>Patient history</td>
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<td>HZ</td>
<td>68 (2.1)</td>
<td>12 (2.1)</td>
<td>4 (1.3)</td>
<td>27 (3.0)</td>
</tr>
<tr>
<td>HZ vaccination</td>
<td>90 (2.8)</td>
<td>15 (2.6)</td>
<td>4 (1.3)</td>
<td>35 (3.9)</td>
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<tr>
<td>VTE</td>
<td>53 (1.7)</td>
<td>9 (1.6)</td>
<td>3 (1.0)</td>
<td>23 (2.5)</td>
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<td>Number of CV risk factors†</td>
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<tr>
<td>0</td>
<td>739 (23.0)</td>
<td>133 (23.0)</td>
<td>82 (26.1)</td>
<td>150 (16.5)</td>
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<tr>
<td>1</td>
<td>1073 (33.4)</td>
<td>202 (34.9)</td>
<td>105 (33.4)</td>
<td>308 (34.0)</td>
</tr>
<tr>
<td>2</td>
<td>883 (27.5)</td>
<td>160 (27.6)</td>
<td>82 (26.1)</td>
<td>251 (27.7)</td>
</tr>
<tr>
<td>3</td>
<td>394 (12.3)</td>
<td>63 (10.9)</td>
<td>36 (11.5)</td>
<td>151 (16.6)</td>
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<tr>
<td>4+</td>
<td>120 (3.7)</td>
<td>21 (3.6)</td>
<td>9 (2.9)</td>
<td>47 (5.2)</td>
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<tr>
<td>Presence of any CV risk factor</td>
<td>2470 (77.0)</td>
<td>446 (77.0)</td>
<td>232 (73.9)</td>
<td>757 (83.5)</td>
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<tr>
<td>Prior CV event</td>
<td>384 (12.0)</td>
<td>62 (10.7)</td>
<td>27 (8.6)</td>
<td>116 (12.8)</td>
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Table 1 Continued

<table>
<thead>
<tr>
<th>Parameter, n (%)</th>
<th>RA 15mg</th>
<th>ADA 40mg</th>
<th>MTX N=314</th>
<th>PsA 15mg</th>
<th>ADA 40mg</th>
<th>MTX N=429</th>
<th>AS 15mg</th>
<th>ASPA N=182</th>
<th>AD 15mg</th>
<th>UPA 30mg</th>
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<tr>
<td></td>
<td>QD N=3209</td>
<td>EOW N=579</td>
<td></td>
<td>QD N=907</td>
<td>EOW N=429</td>
<td></td>
<td>QD N=1182</td>
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<td>QD N=1340</td>
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<td>History of hypertension</td>
<td>1301 (40.5)</td>
<td>252 (43.5)</td>
<td>112 (35.7)</td>
<td>403 (44.4)</td>
<td>179 (41.7)</td>
<td>35 (19.2)</td>
<td>149 (11.1)</td>
<td>123 (9.1)</td>
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<tr>
<td>Diabetes mellitus</td>
<td>382 (11.9)</td>
<td>61 (10.5)</td>
<td>31 (9.9)</td>
<td>122 (13.5)</td>
<td>47 (11.0)</td>
<td>8 (4.4)</td>
<td>26 (1.9)</td>
<td>27 (2.0)</td>
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<tr>
<td>Tobacco/nicotine use‡</td>
<td>1221 (38.0)</td>
<td>199 (34.4)</td>
<td>120 (38.2)</td>
<td>385 (42.4)</td>
<td>163 (38.0)</td>
<td>100 (54.9)</td>
<td>418 (31.2)</td>
<td>418 (30.9)</td>
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<tr>
<td>Elevated LDL-C (&gt;3.36 mmol/L)</td>
<td>869 (27.2)</td>
<td>170 (29.4)</td>
<td>86 (27.5)</td>
<td>253 (28.7)</td>
<td>121 (28.7)</td>
<td>43 (23.8)</td>
<td>197 (14.9)</td>
<td>185 (14.0)</td>
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<tr>
<td>Lowered HDL-C (&lt;1.034 mmol/L)</td>
<td>354 (11.0)</td>
<td>53 (9.2)</td>
<td>39 (12.4)</td>
<td>176 (19.6)</td>
<td>95 (22.4)</td>
<td>34 (18.7)</td>
<td>178 (13.4)</td>
<td>168 (12.5)</td>
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</table>

*Measures of disease activity are as follows: RA, DAS28 (CRP); PsA, DAPSA; AS, ASDAS.
†CV risk factors include history of CV event, hypertension, diabetes mellitus, tobacco/nicotine use, elevated LDL-C, lowered HDL-C.
‡Includes current and former use.

AD, atopic dermatitis; ADA, adalimumab; AS, ankylosing spondylitis; ASDAS, Ankylosing Spondylitis Disease Activity Score; BMI, body mass index; CRP, C reactive protein; csDMARD, conventional synthetic disease-modifying antirheumatic drug; CV, cardiovascular; DAPSA, Disease Activity index for Psoriatic Arthritis; DAS28, Disease Activity Score in 28 Joints; EASI, Eczema Area and Severity Index; EOW, every other week; HDL-C, high-density lipoprotein cholesterol; HZ, herpes zoster; LDL-C, low-density lipoprotein cholesterol; MTX, methotrexate; N/A, not applicable; NSAID, non-steroidal anti-inflammatory drug; PsA, psoriatic arthritis; QD, once a day; RA, rheumatoid arthritis; UPA, upadacitinib; VTE, venous thromboembolism.
Table 2  Exposure and overview of TEAEs

<table>
<thead>
<tr>
<th>Parameter</th>
<th>RA UPA 15 mg QD N=3209</th>
<th>ADA 40 mg EOW N=579</th>
<th>MTX N=314</th>
<th>PsA UPA 15 mg QD N=907</th>
<th>ADA 40 mg EOW N=429</th>
<th>AS UPA 15 mg QD N=182</th>
<th>AD UPA 15 mg QD N=1340</th>
<th>UPA 30 mg QD N=1353</th>
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<tr>
<td><strong>Exposure</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total (PY)</td>
<td>9079.1</td>
<td>1307.7</td>
<td>781.7</td>
<td>1872.3</td>
<td>903.7</td>
<td>320.1</td>
<td>2035.8</td>
<td>2118.0</td>
</tr>
<tr>
<td>Median (minimum, maximum) (years)*</td>
<td>3.46 (0, 5.45)</td>
<td>2.23 (0.04, 5.44)</td>
<td>2.57 (0.02, 5.15)</td>
<td>2.25 (0, 3.9)</td>
<td>1.5 (0.04, 3.22)</td>
<td>1.76 (0.02, 3.26)</td>
<td>1.62 (0, 2.75)</td>
<td>1.65 (0, 2.84)</td>
</tr>
<tr>
<td><strong>Overall TEAEs, E/100 PYs (95% CI)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any AE</td>
<td>205.5 (202.5, 208.5)</td>
<td>203.6 (196.0, 211.5)</td>
<td>206.9 (196.9, 217.2)</td>
<td>244.8 (237.8, 252.0)</td>
<td>229.9 (220.2, 240.0)</td>
<td>241.2 (224.5, 258.8)</td>
<td>250.5 (243.7, 257.5)</td>
<td>278.1 (271.0, 285.2)</td>
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<tr>
<td>Any serious AE</td>
<td>12.4 (11.7, 13.2)</td>
<td>13.7 (11.8, 15.8)</td>
<td>9.6 (7.5, 12.0)</td>
<td>11.1 (9.7, 12.7)</td>
<td>9.0 (7.1, 11.1)</td>
<td>6.6 (4.1, 10.0)</td>
<td>7.1 (6.0, 8.4)</td>
<td>8.2 (7.0, 9.5)</td>
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<tr>
<td>Any AE leading to discontinuation</td>
<td>4.9 (4.4, 5.3)</td>
<td>5.9 (4.6, 7.4)</td>
<td>5.8 (4.2, 7.7)</td>
<td>5.4 (4.4, 6.6)</td>
<td>5.5 (4.1, 7.3)</td>
<td>5.3 (3.1, 8.5)</td>
<td>4.5 (3.6, 5.5)</td>
<td>5.3 (4.4, 6.4)</td>
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<tr>
<td>Deaths,† E/100 PY (95% CI)</td>
<td>0.8 (0.6, 1.0)</td>
<td>0.9 (0.5, 1.6)</td>
<td>0.8 (0.3, 1.7)</td>
<td>0.8 (0.4, 1.3)</td>
<td>0.1 (0.0, 0.6)</td>
<td>0</td>
<td>0</td>
<td>0.1 (0.0, 0.4)</td>
</tr>
</tbody>
</table>

*Minimum in days are as follows: RA: UPA 15 mg, 2 days; ADA 40 mg, 14 days; MTX, 7 days. PsA: UPA 15 mg, 1 day; ADA 40 mg, 14 days. AS: UPA 15 mg, 6 days. AD: UPA 15 mg, 1 day; UPA 30 mg, 1 day.
†Non-treatment emergent deaths included.
AD, atopic dermatitis; ADA, adalimumab; AE, adverse event; AS, ankylosing spondylitis; EOW, every other week; MTX, methotrexate; PsA, psoriatic arthritis; E/100 PY, exposure-adjusted rates per 100 patient years; PY, patient-years; QD, once a day; RA, rheumatoid arthritis; TEAE, treatment-emergent adverse event; UPA, upadacitinib.
Figure 1  Exposure-adjusted event rates for TEAEs of special interest. †Defined as cardiovascular death, non-fatal myocardial infarction and non-fatal stroke. §Including deep vein thrombosis and pulmonary embolism. *RA: UPA 15 mg QD (n=3209), ADA 40 mg EOW (n=579), MTX (n=314); PsA: UPA 15 mg QD (n=907), ADA 40 mg EOW (n=429); AS: UPA 15 mg QD (n=182); AD: UPA 15 mg QD (n=1340), UPA 30 mg QD (n=1353). AD, atopic dermatitis; ADA, adalimumab; AS, ankylosing spondylitis; CPK, creatine phosphokinase; E, event; EOW, every other week; GI, gastrointestinal; MACE, major adverse cardiovascular event; MTX, methotrexate; NMSC, non-melanoma skin cancer; PsA, psoriatic arthritis; PY, patient years; QD, once a day; RA, rheumatoid arthritis; TB, tuberculosis; TEAE, treatment-emergent adverse event; UPA, upadacitinib; VTE, venous thromboembolic event.
NMSC over time of exposure to upadacitinib across diseases and doses (online supplemental figure S4). No pattern or cluster of malignancies excluding NMSC was observed in RA, PsA and AD (online supplemental table S7). Only one malignancy excluding NMSC of stage IVa squamous cell carcinoma of the tongue was observed in the AS study; the patient was a former smoker with less than 5 months of drug exposure. The age–gender adjusted SIR for malignancies other than NMSC using malignancy data from the SEER 18 Registry Research Data 2000–2018 for the general population was estimated at 1.00 (95% CI 0.80 to 1.24) for the upadacitinib 15 mg group in RA, 0.81 (95% CI 0.41 to 1.46) in PsA and 0.50 (95% CI 0.10 to 1.47) in AD; the ratio for patients receiving upadacitinib 30 mg in AD was 1.10 (95% CI 0.53 to 2.03).

Non-melanoma skin cancer
Rates of NMSC (≤0.8 E/100 PY) were generally consistent across diseases for upadacitinib with no events observed in AS. However, event rates of NMSC were numerically higher with upadacitinib compared with adalimumab and no events were reported with MTX. Slightly higher rates of NMSC were also observed in the upadacitinib 30 mg group vs the 15 mg group in AD. Events of NMSC were generally non-serious and did not lead to treatment discontinuation.

Major adverse cardiovascular event
MACE was reported across all treatment groups with rates of ≤0.5/100 PY, and no events were observed with upadacitinib 15 mg in AS. Rates of MACE were comparable between upadacitinib, adalimumab and MTX in RA and PsA (0.3 E/100 PY–0.4 E/100 PY). One event of MACE (<0.1 E/100 PY) was reported each in the upadacitinib 15 mg and 30 mg groups of the AD programme. There was no observed relationship between time of exposure to upadacitinib and MACE (online supplemental figure S5). Most patients experiencing MACE had at least one CV risk factor. There were 11 CV deaths, all within the RA population.

Venous thromboembolism
VTE was observed among patients receiving upadacitinib across disease states with rates of 0.4 E/100 PY in RA, 0.2 E/100 PY in PsA, 0.3 E/100 PY in AS, and <0.1 E/100 PY for both upadacitinib 15 mg and 30 mg in AD. Rates with upadacitinib were comparable to those observed with adalimumab (RA and PsA) and MTX (RA). There was no relationship observed between time of exposure to upadacitinib and VTE (online supplemental figure S6). Events of PE were more frequent than DVT in RA, PsA and AS; in AD, PE and/or DVT, rates were <0.1 E/100 PY each for upadacitinib 15 mg and upadacitinib 30 mg. Most patients experiencing a VTE had at least one CV and/or thromboembolic risk factor. There were two fatal VTE events across all diseases, both being PEs in patients with RA.

Laboratory abnormalities
AEs of elevations in blood CPK levels occurred in all diseases, were dose dependent, occurred more frequently with upadacitinib than with adalimumab and MTX and were primarily asymptomatic. Few patients had symptoms of muscle pain, and alternative aetiologies of vigorous physical activity were identified in most cases. One case of rhabdomyolysis occurred in an adolescent patient receiving upadacitinib 30 mg for AD following a boxing activity. AEs of anaemia, neutropenia and lymphopenia were also reported in RA, PsA, AS and AD. Most laboratory abnormalities were reported to be mild overall and transient and few resulted in discontinuation of the study drug (these abnormalities are further discussed in the online supplemental material).

AEs related to laboratory parameters of special interest for the JAK inhibitor class (anaemia, lymphopenia, neutropenia, hepatic disorders and CPK elevations) are further discussed in the online supplemental material 1; abnormal laboratory results of haematology and chemistry tests by grading criteria performed during the upadacitinib studies are described in online supplemental tables S8 and S9.

AESIs by age
While numbers of patients aged ≥65 years are limited overall, event rates tended to be higher for MACE, VTE, malignancies and serious infections, regardless of treatment (upadacitinib, adalimumab or MTX; online supplemental table S10). In RA and PsA, rates of these events in patients aged ≥65 years were similar between upadacitinib 15 mg and adalimumab.

Additional information on AESIs by concomitant therapy
Overall, the event rates of AESIs occurring in patients receiving upadacitinib monotherapy or upadacitinib combination therapy with a csDMARD in RA and PsA were generally consistent (online supplemental figure S7).

DISCUSSION
This safety analysis of 6991 patients receiving upadacitinib with a maximum of 5.45 years of follow-up appears largely consistent across RA, PsA, AS and AD for key AESIs such as MACE, VTE and malignancy excluding NMSC, and no new or unexpected safety risks were identified compared with previous reports.24–27 Differences in AE types and rates across disease states reflect expected distinctions between disease-specific patient populations and background risks.28–31 Long-term exposures with adalimumab (RA and PsA) and MTX (RA only), although with lower numbers of patients evaluated, within the upadacitinib clinical trial programme further enabled the contextualisation of the described upadacitinib safety profile alongside active comparators.

Varied comorbidities and patient populations across diseases likely resulted in differences in some TEAE types and rates.28–31 Notably, acne was one of the most common
TEAEs within the AD population while minimally represented across others, possibly owing to a younger overall population of patients with AD or greater scrutiny of dermatological AEs by dermatologists. Acne has been observed in all trials of JAK inhibitors in AD, though most cases are mild to moderate and do not result in treatment discontinuation. \(^5\)–\(^7\) Additionally, baseline concomitant corticosteroid use was higher among patients with RA, largely reflecting differences in treatment recommendations and guidelines; however, the relationship between concomitant corticosteroid use and events of MACE or VTE have not yet been explored. \(^8\)–\(^15\) Determining a relationship between corticosteroid use and MACE or VTE is beyond the scope of this analysis, given the challenge of separating corticosteroid use from other risk factors, the need for further data regarding dosing (mean, median and cumulative) and given the included data concerns corticosteroid use at baseline only. The event rates of herpes zoster, NMSC and elevation in CPK levels were higher with upadacitinib compared with adalimumab and MTX in RA and/or PsA; however, the rates of the remaining AESIs varied without consistent findings between upadacitinib vs the active comparators in the two programmes. An increased risk of herpes zoster and elevated CPK levels is associated with JAK inhibition and consistent with the overall safety profiles of JAK inhibitors. \(^8\)–\(^24\) \(^36\)–\(^37\) Herpes zoster infections observed with upadacitinib were largely non-serious and limited to one dermatome, while elevations in CPK levels were mostly asymptomatic and transient. Of note, the upadacitinib label recommends herpes zoster vaccination for all patients prior to initiating upadacitinib therapy. Furthermore, there is a recognised potential for JAK inhibitors to be associated with an increased risk of NMSC, with systematic reviews of the use of tofacitinib, ruxolitinib and baricitinib identifying higher rates of NMSC in patients with IMIDs. \(^38\)–\(^40\) Rates of NMSC were higher in patients receiving upadacitinib than active comparators, primarily presenting as basal or squamous cell carcinoma in patients aged ≥65 years. Further studies with more long-term data are required to fully determine any association between JAK inhibitors and NMSC. AESIs were similar across all diseases, except for an increase in opportunistic infections excluding herpes zoster in AD due to eczema herpeticum, which is commonly associated with AD. \(^41\) An overall increased risk of TEAEs was observed in the upadacitinib 30 mg population compared with upadacitinib 15 mg for AD. COVID-19 infections had a notable impact on serious infection rates, especially in PsA. The timing of the PsA studies coinciding with that of the COVID-19 pandemic likely contributed, and when COVID-19 infections were excluded, there was little variation in serious infections with upadacitinib across diseases.

Recent data from ORAL Surveillance, an event-driven, head-to-head study in an RA population enriched for CV risk, demonstrated that tofacitinib failed to meet the non-inferiority criteria for events of MACE and malignancy (excluding NMSC) vs TNF inhibitor therapy. \(^10\) Patients most at risk of experiencing events of MACE or malignancy while receiving tofacitinib compared with TNF inhibitors were aged ≥65 years and had a history of smoking, underscoring the need to evaluate patient medical history when selecting therapeutic options. \(^10\) \(^42\) Of note, a similar signal to that observed in ORAL Surveillance has not been observed in the overall tofacitinib trial programme, and the separation between tofacitinib and TNF inhibitors only became apparent in a patient population enriched for overall risk. A definitive safety outcomes study like ORAL Surveillance has not been conducted with upadacitinib and whether the potential increased risk of these events reflects a JAK inhibitor class effect remains unclear.

Events of malignancies excluding NMSC were reported in patients receiving upadacitinib and were consistent with active comparators, with age-adjusted SIR data suggestive of no increased risk of these events compared with the general population. Numerically higher rates of malignancy were observed with upadacitinib 30 mg compared with upadacitinib 15 mg in AD; however, four of the nine malignancies observed with upadacitinib 30 mg occurred within 6 months after initiating upadacitinib.

VTE risk with JAK inhibitor therapy has become a concern owing to elevated rates observed with 4 mg baricitinib compared with placebo in several studies. \(^6\)–\(^7\) A higher rate of VTE was also observed with tofacitinib 10 mg given two times a day compared with TNF inhibitors in ORAL Surveillance. \(^10\) Although events of VTE were reported in patients receiving upadacitinib across diseases, the rates appeared consistent with those observed for adalimumab in RA and PsA and MTX in RA and were consistent with background rates for the individual diseases. \(^43\)–\(^44\) Several risk factors for VTE are inherently present in IMIDs including systemic inflammation, and patients with RA are at a substantially increased risk of VTE compared with the general population. \(^44\)–\(^46\) The potential association between JAK inhibitors and VTE risk requires further analysis, given the inherent risk carried by the conditions for which JAK inhibitors are indicated.

Limitations of this analysis include a lack of extended placebo control arms for comparative analysis and lack of postmarketing data. Additionally, the pooling strategy for upadacitinib leveraged data sets which did not include active comparator arms. Substantial exposure time is represented; however, data for certain populations (AS) and active comparators are more limited and occurrence of certain events was low. Consequently, rate estimations for events such as malignancy, MACE and VTE have limited precision for conclusive interpretations. Additional follow-up is required to fully characterise the rates observed on upadacitinib for these and other long latency events. All reported TEAEs were reported in the closely monitored clinical trial setting, and the full safety profile of upadacitinib should also consider reported TEAEs from clinical practice settings and registries. Finally, only indications with approval as of December 2021 were included in this analysis, and, as such, UC was
not included, thus limiting the further characterisations of events that may have a dose response.

CONCLUSIONS
Analyses of long-term data demonstrate that upadacitinib was generally well tolerated in RA, PsA, AS and AD with no new safety risks identified compared with previous reports. Some variations in events were observed across diseases, possibly reflecting differences in patient populations and disease-associated comorbidities that impact background risk. Follow-up of patients receiving upadacitinib will continue as these trials are ongoing.

Author affiliations
1. Department of Rheumatology and Clinical Immunology, Charité Universitätsmedizin Berlin, Berlin, Germany
2. Department of Rheumatology, Metroplex Clinical Research Center, Dallas, Texas, USA
3. Division of Infectious Diseases, Oregon Health and Science University, Portland, Oregon, USA
4. School of Medicine, Griffith University School of Medicine, Brisbane, Queensland, Australia
5. Department of Clinical Medicine, Trinity College Dublin, Dublin, Ireland
6. Wellcome-HRB Clinical Research Facility, St. James’ Hospital, Dublin, Ireland
7. Rheumatology, Organización Medica de Investigacion, Buenos Aires, Argentina
8. The First Department of Internal Medicine, University of Occupational and Environmental Health Japan, Kitakyushu, Japan
9. AbbVie Inc, North Chicago, Illinois, USA
10. AbbVie Ltd, Maidenhead, UK
11. Rheumatology Research Division, Swedish Medical Center/Providence St. Joseph Health, Seattle, Washington, USA
12. Department of Dermatology and Laboratory for Inflammatory Skin Diseases, Icahn School of Medicine at Mount Sinai, New York, New York, USA

Twitter Peter Nash @drpnash
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Patient consent for publication Not applicable.

Ethics approval This study involves human participants and is an integrated analysis of a clinical trials programme. All trials were conducted according to the International Conference on Harmonisation guidelines, the Declaration of Helsinki principles and applicable local country regulations. All study-related documents were approved by independent ethics committees and institutional review boards at each site. All patients provided written informed consent.

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Data availability statement Data are available upon reasonable request. AbbVie is committed to responsible data sharing regarding the clinical trials we sponsor. This includes access to anonymized individual and trial-level data (analysis data sets), as well as other information (eg, protocols, clinical study reports, or analysis plans), as long as the trials are not part of an ongoing or planned regulatory submission. This includes requests for clinical trial data for unlicensed products and indications. These clinical trial data can be requested by any qualified researchers who engage in rigorous, independent, scientific research, and will be provided following review and approval of a research proposal, Statistical Analysis Plan (SAP), and execution of a Data Sharing Agreement (DSA). Data requests can be submitted at any time after approval in the US and Europe and after acceptance of this manuscript for publication. The data will be accessible for 12 months, with possible extensions considered. For more information on the process or to submit a request, visit the following link: https://www.abbvieclinicaltrials.com/hcp/data-sharing/.

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ORCID iDs
Gerd R Burmester http://orcid.org/0000-0001-7518-1131
Stanley B Cohen http://orcid.org/0000-0002-9226-679X
Kevin L Winthrop http://orcid.org/0000-0002-3892-6947
Peter Nash http://orcid.org/0000-0002-2571-788X
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