Liver Injury with Ulipristal Acetate: Exploring the Underlying Pharmacological Basis

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Abstract

Introduction The European Medicines Agency has suspended the use of ulipristal acetate (UPA) in the treatment of uterine fibroids and is reassessing its association with a risk of liver injury.

Objectives Our objectives were to characterize the post-marketing reporting of drug-induced liver injury (DILI) with UPA and investigate the underlying pharmacological basis.

Methods We queried the worldwide FDA Adverse Event Reporting System and performed a disproportionality analysis, selecting only hepatic designated medical events (DMEs) where UPA was reported as suspect. The reporting odds ratios (RORs) were calculated, and we considered a lower limit of the 95% confidence interval (LL95% CI) > 1 as significant. Physiochemical/pharmacokinetic features were extracted to assess the risk of hepatotoxicity by applying predictive DILI risk models. Mifepristone and leuprolide were selected as comparators.

Results A significantly higher proportion of liver disorders was reported for UPA than for mifepristone (2.9 vs. 0.8%; \( p < 0.00001 \)) and leuprolide (2.9 vs. 1.6%; \( p = 0.015 \)). As regards hepatic DMEs, statistically significant RORs were found for autoimmune hepatitis (\( N = 5; \) LL95% CI 16.8), DILI (\( n = 5; \) LL95% CI 5.9), and acute hepatic failure (\( N = 5; \) LL95% CI 9.3). No signals of DILI emerged for mifepristone and leuprolide acetate. UPA and mifepristone showed high lipophilicity and hepatic metabolism (predicted intermediate DILI risk). Leuprolide exhibited contrasting features, resulting in no DILI concern. Inhibition of different liver transporters and the presence of a reactive metabolite were also recognised for UPA.

Conclusion Different drug properties previously linked to the occurrence of DILI may partially explain the reporting pattern observed with UPA. Our “bedside-to-bench” approach may support regulators in the risk–benefit assessment of UPA.

1 Introduction

Ulipristal acetate (UPA) is an orally active selective progesterone-receptor modulator. In 2012, the European Medicines Agency (EMA) approved UPA 5 mg/day for the treatment of moderate to severe symptoms of uterine fibroids in adult women of reproductive age, with the treatment duration limited to 3 months [1]. It is also approved for emergency contraception as one single 30 mg dose.

In February 2018, the EMA issued temporary restrictive measures for UPA after five cases of drug-induced liver injury (DILI) were reported, four of which required liver transplantation. The Pharmacovigilance Risk Assessment Committee (PRAC) subsequently made temporary recommendations advising physicians not to take on new patients or initiate new treatment courses. In May 2018, the status of UPA as a potential DILI-inducing agent was neither confirmed nor fully ruled out; however, in eight cases of serious liver injury the role of UPA was deemed possible, and the PRAC made recommendations to minimize the risk of liver injury by forbidding its use in women with underlying hepatic disorders or liver enzyme levels more than twice the upper limit of normal [2–4].

Subsequent analyses reported that elevation in liver enzyme levels was recognised in 11 of 1868 women (0.6%) treated with UPA in pivotal trials, of which only one received the approved dosage of 5 mg/day, and none met Hy’s Law [5, 6]. Furthermore, a retrospective study assessing liver
Key points

This original research explored the possible underlying pharmacological basis involved in liver injury associated with ulipristal acetate by applying a “bedside-to-bench” mixed approach that included estimating the extent of life-threatening hepatic reactions in the real world and assessing physiochemical and pharmacokinetic parameters correlated with drug-induced liver injury.

The over-reporting of severe life-threatening hepatic reactions, particularly autoimmune hepatitis, associated with ulipristal acetate in post-marketing surveillance may be partially explained by its physiochemical (high lipophilicity) and pharmacokinetic (hepatic metabolism, long half-life, inhibition of liver transporters, reactive metabolite formation) features.

This mixed approach may be helpful for regulators in supporting risk-benefit evaluation during the current suspension issued by the European Medicines Agency for ulipristal acetate in the management of uterine fibroids.

3 Methods

3.1 Pharmacovigilance Evaluation

We performed an observational retrospective disproportionality analysis to highlight and characterize hepatic AEs of clinical interest with (higher than expected) increased reporting. The FAERS database (public dashboard), the US repository of AEs and medication errors comprising more than 18 million reports gathered worldwide, was queried to retrieve UPA reports recorded between the first quarter of 2011 and the fourth quarter of 2019 (search performed on 8 April 2020). To assign a clinical priority to emerging hepatic safety issues, we used the public list of DMEs developed by the EMA [9]. Specifically, seven of the 62 different reactions included in the public list of DMEs concern hepatic AEs (acute hepatic failure, autoimmune hepatitis, DILI, hepatic failure, hepatic infarction, hepatic necrosis, and hepatitis fulminant). The reporting odds ratio (ROR) with relevant 95% confidence interval (CI) was calculated as a measure of disproportionality, using all other drugs/events recorded in FAERS as a comparator. Traditional criteria for signal detections were used, i.e., lower limit of the 95% CI of the ROR > 1 with at least three cases of interest reported [10]. Cases were manually checked for duplicates using case identification and overlapping information among records, and, for ROR calculation, a case counts as many-fold as the number of hepatic events reported. Case-by-case assessment for concomitant drugs with hepatotoxic potential was also performed based on classification proposed by Björnsson et al. [11], focusing on agents in category A and B.

For comparison, we also extracted data on mifepristone, a well-known progesterone-receptor antagonist available for pregnancy termination [12], and leuprolide acetate, a synthetic agonist analogue of gonadotropin-releasing hormone first approved for the treatment of advanced prostatic cancer and also currently used for the management of uterine fibroids [13]. Mifepristone and leuprolide acetate were, respectively, selected according to affinity of chemical structure and comparable therapeutic indication (i.e., uterine fibroids).

2 Objective

In this “regulatory limbo”, pharmacology may aid investigations into the potential underlying mechanisms of DILI associated with UPA and with estimating the extent of reporting in post-marketing surveillance.

Therefore, to explore the relationship between the use of UPA and the occurrence of DILI, we implemented a “two-step” approach based on (1) analysis of spontaneous reports submitted to the US FDA Adverse Event Reporting System (FAERS) database, including disproportionality analysis for hepatic reactions classified as designated medical events (DMEs), namely rare serious adverse events (AEs) with a recognized drug-attributable risk [9]; and (2) review of physiochemical and pharmacokinetic features known to be potentially involved in DILI.
metabolism, and inhibition of hepatic transporters can confer a risk for DILI, which may increase susceptibility to hepatotoxicity because of host-related factors such as female sex and age [14, 15].

We also queried public online prediction tools (namely, ADVERPred [16] and Vienna LiverTox Workspace [17]) to investigate the existence of physiochemical features associated with DILI and the interaction profile with liver transporters potentially involved in hepatic damage. Additionally, we applied the DILI risk model proposed by Chen et al. [18] based on lipophilicity, formation of reactive metabolites, and scheduled dose/peak concentration (C_max). According to the model, a score > 7 indicates the most DILI concern/solid evidence, a score of 3–7 indicates less DILI concern/weak evidence, and a score < 3 indicates no DILI concern/no evidence of hepatotoxicity. This so-called “rule-of-two” (Ro2) model, validated using 354 drugs, is a simple way to predict the severity of DILI in humans and has also been successfully applied to stratify DILI liability among direct-acting antivirals for hepatitis C and cyclin-dependent kinase 4/6 inhibitors to treat hormone receptor-positive breast cancer [14, 19].

Finally, we also reviewed the physiochemical and pharmacokinetic features of mifepristone and leuprolide acetate as a comparison.

4 Results

Data collected from FAERS for UPA, mifepristone, and leuprolide acetate are shown in Table 1. Overall, 578, 6220, and 44,079 reports mentioning UPA, mifepristone, and leuprolide acetate, respectively, as the suspect agent were found. Serious cases ranged from 51.1% for mifepristone to 92.0% for UPA. A significantly higher proportion of liver disorders was reported for UPA than for mifepristone (2.9 vs. 0.8%; p < 0.00001) and leuprolide acetate (2.9 vs. 1.6%; p = 0.015) in the real world, although notoriety bias (i.e., increased reporting of hepatic injuries following increased awareness of submitters) cannot be ruled out, given that eight of 17 reports were recorded after warnings issued by the EMA. As regards hepatic DMEs reported with UPA, statistically significant RORs were found for autoimmune hepatitis (N = 5; ROR 40.6; 95% CI 16.8–97.9), DILI (N = 5; ROR 14.2; 95% CI 5.9–34.3), and acute hepatic failure (N = 5; ROR 22.5; 95% CI 9.3–54.2).

Conversely, no significant ROR was found for hepatic DMEs reported with mifepristone (eight cases of hepatic failure and three of DILI) and leuprolide acetate (Supplementary Materials 1). Overall, 60 hepatic DMEs were reported with leuprolide acetate, with hepatic failure (N = 32) the most represented. However, these events occurred in patients affected by advanced prostatic or breast cancer, and no life-threatening hepatic events were reported in the 17 women treated for uterine fibroids. Notably, concomitant hepatotoxic agents were retrieved in only 17.6% of UPA reports (all in category A) and in none of the autoimmune hepatitis cases.

The physiochemical and pharmacokinetic parameters retrieved for UPA [20, 21], mifepristone [22, 23], and leuprolide acetate [13, 24] are summarized in Table 2. Both UPA and mifepristone exhibited high lipophilicity (respectively, LogP 4.45 and 5.43), hepatic metabolism, and inhibition of different liver transporters. A reactive metabolite was found for UPA, and Ro2 was positive for mifepristone. Prediction tools showed no risk of hepatotoxicity for these agents, except for cholestasis potentially due to mifepristone. Implementation of dose- and C_max-based DILI scores resulted in less DILI concern/weak evidence of hepatotoxicity for both UPA (respectively, 4.82 and 5.75 patients) and mifepristone (respectively, 4.43 and 4.40 patients).

Conversely, leuprolide acetate showed contrasting properties, namely poor lipophilicity (LogP 1.04), lack of hepatic metabolism, and no impact on liver transporters. Prediction tools reported no risk of hepatotoxicity, and dose- and C_max-based DILI scores resulted in no DILI concern/no evidence of hepatotoxicity (respectively, 0.24 and 0.31 patients).

5 Discussion

Our “two-step” analysis showed that the reporting patterns for life-threatening hepatic AEs were quite different among the drugs of interest, with higher reporting of autoimmune hepatitis and DILI with UPA, whereas similar physiochemical features were shared with mifepristone (Table 2), indicating that investigation of underlying pharmacological mechanisms is required, especially exploring the immune-mediated hypothesis.

Notably, specific pharmacological properties of UPA may explain the non-negligible reporting of certain hepatic DMEs. In particular, a minor reactive, partially characterized metabolite was detected in human faeces, proposed to be a glutathione conjugate of mono-oxygenated UPA. This proposed structure is consistent with the oxidation of the 4,5 carbon atoms to a reactive epoxide, followed by deactivation through glutathione conjugation [25]. This reactive metabolite may covalently bind to a protein and form a hapten–protein adduct and thereby elicit an adaptive immune response in susceptible individuals, potentially leading to occurrence of autoimmune hepatitis and DILI. Furthermore, this phenomenon may be enhanced by high lipophilicity and inhibition of breast cancer resistance protein (BCRP) in liver cells as well as predicted bile salt export pump (BSEP) inhibition in silico by UPA. Finally, dose dependency may also play a

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Table 1  Summary of spontaneous reports retrieved from the FAERS database concerning ulipristal acetate and mifepristone

| FAERS parameters | Ulipristal acetate | Mifepristone | Leuprolide acetate |
|------------------|-------------------|--------------|-------------------|
| Total reports    | 578               | 6220         | 44,079            |
| Age distribution, years |                |              |                   |
| <18              | 20 (3.4)         | 116 (1.9)    | 1609 (3.7)        |
| 18–64            | 364 (63.0)       | 4686 (75.3)  | 12,414 (28.2)     |
| 65–85            | 0 (0.0)          | 123 (2.0)    | 10,021 (22.7)     |
| >85              | 0 (0.0)          | 3 (0.1)      | 1713 (3.9)        |
| Not specified    | 194 (33.6)       | 1292 (20.7)  | 18,322 (41.5)     |
| Serious case     | 532 (92.0)       | 3177 (51.1)  | 27,258 (61.8)     |
| Proportion of deaths | 5 (0.9)       | 274 (4.4)    | 6860 (15.6)       |
| Most frequent AEs | Unintended pregnancy | Haemorrhage | Hot flush |
|                  | 159 (27.5)       | 1748 (28.1)  | 4203 (9.5)        |
| Abortion spontaneous | 107 (18.5)    | 1714 (27.6)  | 3997 (9.1)        |
| Vaginal haemorrhage | 64 (11.1)     | 766 (12.3)   | 2185 (5.0)        |
| Drug ineffective | 44 (7.6)         | 586 (9.4)    | 2074 (4.7)        |
| Pregnancy after post-coital contraception | 493 (7.9) |            | 1757 (4.0)        |
| Hepatobiliary disorders | 17 (2.9) | 48 (0.8) | 722 (1.6) |
| Mean age         | 36.6 ± 9.0       | 40.5 ± 16.5  | 63.0 ± 19.1       |
| Sex              | 16 F; 1 NA       | 25 F; 22 NA; 1 M | 218 F; 33 NA; 471 M |
| Serious          | 17 (100.0)       | 45 (93.8)    | 677 (93.8)        |
| Proportion of deaths | 0 (0.0)      | 3 (6.3)      | 129 (17.9)        |
| Reporter country |                  |              |                   |
| Germany          | 6                | –            | 9                 |
| Italy            | 3                | –            | 13                |
| Spain            | 3                | –            | 10                |
| France           | 2                | 2            | 40                |
| UK               | 1                | –            | 7                 |
| Portugal         | 1                | –            | 2                 |
| Hungary          | 1                | –            | 3                 |
| USA              | –                | 32           | 119               |
| Canada           | –                | 1            | 45                |
| Saudi Arabia     | –                | 1            |                   |
| Japan            | –                | –            | 115               |
| Other            | –                | –            | 60                |
| Not specified    | –                | 12           | 292               |
| Reason for use   |                  |              |                   |
| Uterine leiomyoma | 11               | –            | 17                |
| Menometrorrhagia | 6                | –            | 5                 |
| Dysmenorrhoea    | 5                | –            | 1                 |
| Post-coital contraception | 3 | – | – |
| Cushing’s syndrome | –     | 35          | –                 |
| Abortion induced | –                | 8            | –                 |
| Placenta accreta | –                | 1            | –                 |
| Endometriosis    | –                | –            | 67                |
| Prostate/breast cancer | – | – | 404 |
| NA               | –                | 4            | 228               |

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Liver Injury Associated with Ulipristal Acetate

Although UPA is used at low dosages for the management of uterine fibroids (one-sixth of the scheduled dosage for emergency contraception), the long-term course of treatment (up to 12 weeks) coupled with the extended half-lives of UPA and its main active metabolite PGL4002 (> 24 h) may lead to dose-dependent accumulation causing hepatotoxicity [20, 27]. Consequently, we fully endorse the PRAC recommendations that encourage strict monitoring of liver function up to 4 weeks after stopping treatment with UPA.

Although prediction tools showed no risk of hepatotoxicity, except for cholestasis potentially due to mifepristone, both UPA and mifepristone exhibited dose- and $C_{\text{max}}$-based DILI scores ranging from 4.4 to 5.75, resulting in less DILI concern/weak evidence (similar to amiodarone, entacapone, moxifloxacin) compared with drugs associated with severe hepatotoxicity risk (i.e., tolcapone, which was withdrawn from the market). These findings are in line with real-world data showing sporadic cases of severe liver injury associated with UPA compared with a post-marketing exposure estimated at > 765,000 patients [5]. Furthermore, cases of cholestatic DILI caused by mifepristone were recently reported [28, 29], supporting predicted data from in silico models.

Conversely, leuprolide acetate exhibits contrasting physicochemical and pharmacokinetic properties (poor lipophilicity, parenteral administration, lack of hepatic metabolism), with low dose- and $C_{\text{max}}$-based DILI scores (< 0.5) resulting in no DILI concern/no evidence of hepatotoxicity. These findings are confirmed by literature data, where leuprolide acetate is classified as an unlikely cause of clinically apparent liver injury, given that only mild serum enzyme elevations in 3–5% of patients were reported [30]. Furthermore, most cases of elevated transaminase levels were not associated with a direct impact of the agent on liver function but were caused by non-alcoholic fatty liver arising from metabolic changes due to an androgen-deprivation state induced by leuprolide acetate.

### Table 1 (continued)

| FAERS parameters                  | Ulipristal acetate | Mifepristone | Leuprolide acetate |
|-----------------------------------|--------------------|--------------|--------------------|
| Hepatic DMEs at PT level (ROR; 95% CI) |                    |              |                    |
| Autoimmune hepatitis              | 5 (40.6; 16.8–97.9) | 1 (NC)       | 3 (0.32; 0.10–0.98) |
| Drug-induced liver injury         | 5 (14.2; 5.9–34.3)  | 3 (0.78; 0.25–2.43) | 7 (0.26; 0.12–0.54) |
| Acute hepatic failure             | 5 (22.5; 9.3–54.2)  | –            | 5 (0.29; 0.15–0.70) |
| Hepatic failure                   | 2 (NC)             | 8 (0.91; 0.45–1.81) | 32 (0.51; 0.36–0.72) |
| Hepatic necrosis                  | –                  | –            | 3 (0.23; 0.07–0.70) |
| Hepatitis fulminant               | –                  | –            | 10 (1.74; 0.94–3.24) |
| Concomitant agents (category)a    |                    |              |                    |
| None                              | 6                  |              | 15                 |
| Pantoprazole (C)                  | 5                  | –            | –                  |
| Metamizole (E)                    | 4                  | –            | –                  |
| Doxycycline (C)                   | 3                  | –            | –                  |
| Decapeptyl (E)                    | 2                  | –            | –                  |
| Desogestrel (A)                   | 2                  | –            | –                  |
| Lansoprazole (C)                  | 1                  | –            | –                  |
| Atorvastatin (A)                  | 1                  | –            | –                  |
| Metoclopramide (E)                | 1                  | –            | –                  |
| Indapamide (E)                    | 1                  | –            | –                  |
| Ethinylestradiol/Levonorgestrel (A)| 1                 | –            | –                  |
| Verapamil (C)                     | –                  | –            | 1                  |
| Triamterene (D)                   | –                  | –            | 1                  |
| Valsartan (D)                     | –                  | –            | 1                  |
| Tramadol (E)                      | –                  | –            | 1                  |

Data are presented as mean ± standard deviation or N (%) unless otherwise indicated

AE adverse event, CI confidence interval, DME designated medical event, F female, FAERS US FDA Adverse Event Reporting System, M male, NA not available, NC not calculated, PT preferred term, ROR reporting odds ratio

aBased on classification reported in Björnsson et al. [11]. Number of convincing reports in the published literature: category A, ≥ 50; B, 12–49; C, 4–11; D, 1–3; E, none

bExtracted from the 17 reports in which leuprolide acetate was used in the management of uterine leiomyoma
Table 2 Summary of physiochemical and pharmacokinetic (PK) features of ulipristal acetate and mifepristone potentially involved in drug-induced liver injury [13, 16–18, 20–25]

| Physiochemical and PK features | Ulipristal acetate | Fulfilled criteria | Mifepristone | Fulfilled criteria | Leuprolide acetate | Fulfilled criteria |
|-------------------------------|-------------------|--------------------|-------------|--------------------|-------------------|--------------------|
| Molecular weight (> 600 Da)   | 475.6             | ✗                  | 429.6       | ✗                  | 1209.4            | ✗                  |
| Scheduled daily dose (≥ 50-100 mg/day)  | 5                | ✗                  | 200         | ✓                  | not available per os | ✗                  |
| Lipophilicity (LogP ≥3)       | 4.45              | ✓                  | 5.33        | ✓                  | 1.04              | ✗                  |
| Topological polar surface area (< 75 Å2) | 63.7             | ✓                  | 40.5        | ✓                  | 429.04            | ✗                  |
| CYt/IC50 ≥ 0.1 BSEP IC50     | No data           | ?                  | No data     | ?                  | No data           | ?                  |
| Oxidative stress             |                   |                    |             |                    |                   |                    |
| Reactive metabolites formation | A reactive metabolite was retrieved in faeces* | ✓              | No          | ✗                  | No                | ✗                  |
| “Rule-of-two”                |                   |                    |             |                    |                   |                    |
| Mitochondrial liability      |                   |                    |             |                    |                   |                    |
| Mitochondrial dysfunction    | No data           | ?                  | No data     | ?                  | No data           | ?                  |
| Metabolism                   |                   |                    |             |                    |                   |                    |
| Hepatic metabolism           | CYP 3A4 +++       | ✓                  | CYP 3A4 +++ | ✓                  | No                | ✗                  |
| Hepatic transporters         |                   |                    |             |                    |                   |                    |
| BSEP inhibition              |                   |                    |             |                    |                   |                    |
| Positive [1.00] (predicted in silico) | ✗               | ✓                  | Yes [1.00] (predicted in silico) | ✓                  | Pos [1.00] (in vitro) | ✓                  |
| MRP 2 transport              |                   |                    |             |                    |                   |                    |
| Positive [0.60] (predicted in silico) | ?                | ✓                  | No data [0.40] (predicted in silico) | ✗                  | Neg [0.00] (in vitro) | ✗                  |
| MRP 3 inhibition             |                   |                    |             |                    |                   |                    |
| Positive [1.00] (predicted in silico) | ?                | ✓                  | No data [1.00] (predicted in silico) | ✗                  | Pos [1.00] (in vitro) | ✓                  |
| MRP 4 inhibition             |                   |                    |             |                    |                   |                    |
| Positive [1.00] (predicted in silico) | ?                | ✓                  | No data [1.00] (predicted in silico) | ✗                  | Pos [1.00] (in vitro) | ✓                  |
| P-glycoprotein inhibition    |                   |                    |             |                    |                   |                    |
| Positive [0.66] (predicted in silico) | ?                | ✓                  | Yes [0.91] (predicted in silico) | ✓                  | Pos [0.97] (in vitro) | ✓                  |
| BCRP inhibition              |                   |                    |             |                    |                   |                    |
| Positive [0.21] (predicted in silico) | ?                | ✓                  | Yes [0.69] (predicted in silico) | ✓                  | Pos [0.13] (in vitro) | ✗                  |
| OATP1B1 inhibition           |                   |                    |             |                    |                   |                    |
| Positive [1.00] (predicted in silico) | ?                | ✓                  | Yes [1.00] (predicted in silico) | ✓                  | Pos [1.00] (in vitro) | ✓                  |
| OATP1B3 inhibition           |                   |                    |             |                    |                   |                    |
| Positive [1.00] (predicted in silico) | ?                | ✓                  | Yes [1.00] (predicted in silico) | ✓                  | Pos [1.00] (in vitro) | ✓                  |
| DILI risk score and prediction |                   |                    |             |                    |                   |                    |
| ADVERPred                    |                   |                    |             |                    |                   |                    |
| Vienna LiverTox Workspace**  | Negative          | ✗                  | Negative    | ✗                  | Negative          | ✗                  |
| Drug-induced liver injury    | 0.30              | ✗                  | 0.12        | ✗                  | 0.59              | ✗                  |
| Hyperbilirubinemia           | 0.27              | ✗                  | 0.02        | ✗                  | 0.24              | ✗                  |
| Cholestasis                  | 0.03              | ✗                  | 1.00        | ✗                  | 0.75              | ✗                  |
| Dose-based DILI score***     | 4.82              | less-DILI-concern | 4.43        | less-DILI-concern | 0.24              | no-DILI-concern    |
| Cmax-based DILI score***     | 5.75              | less-DILI-concern | 4.40        | less-DILI-concern | 0.31              | no-DILI-concern    |

BCRP breast cancer resistance protein, BSEP bile salt export pump, Cmax peak concentration, Cplasma serum concentration, CYP cytochrome P450, DILI drug-induced liver injury, IC50 half-maximal inhibition, MRP multidrug resistance-associated protein, NA not available, Neg negative, OATP1B organic anion-transporting polypeptide, Pos positive

+++ extensive metabolism by CYP450, ++ moderate metabolism by CYP450, + low metabolism by CYP450

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From a clinical perspective, the EMA’s current suspension of marketing authorization for UPA may pose major issues in daily practice, as the efficacy of UPA in reducing fibroid-associated bleeding, duration of surgery, frequency of blood transfusions, and fibroid size when used in the preoperative treatment of uterine fibroids is recognized [31–35]. Additionally, current evidence suggests that prolonged use of UPA may be a good alternative to other more invasive treatment modalities, particularly for clinical scenarios such as bridging the gap until menopause or in the younger patient to minimize symptoms until pregnancy is desired [32]. In this scenario, leuprolide acetate should be considered a potential alternative, given its proven efficacy in reducing fibroid size, intraoperative blood loss, and the frequency of blood transfusion [32–35]; however, clinical trials have reported a significantly higher occurrence of hot flushes compared with UPA, and our findings confirm this [33]. Although no signals of DILI emerged for leuprolide acetate in our pharmacovigilance analysis, a non-negligible number of serious AEs was reported, mostly in different clinical scenarios with respect to UPA (namely older patients affected by advanced prostatic or breast cancer).

As a whole, the hepatotoxic liability of UPA (and its potential definite withdrawal) must be balanced against two main clinical aspects: (1) a potentially significant increase in the number of minor and major surgeries for uterine fibroids, as already seen after the first suspension in 2018 [36], with possible increases in postoperative complications; and (2) the risk/benefit profile of medical alternatives (leuprolide acetate) in the light of real-life findings.

We acknowledge the limitations of this study, related to both pharmacovigilance analyses, including FAERS data (e.g., quality of reports, potential existence of remaining duplicates, reporting biases, lack of exposure data, inability to establish firm causality between drug exposure and occurrence of AEs, and limited verification of events through clinical features) and pharmacokinetic/pharmacodynamic assessment. We selected three agents with different markets, indications, treatment durations, and mechanisms of action, which meant comparison was difficult, particularly given the lack of a clear DILI effect with these drugs. Our study was intended to be a proof of concept, and the design or validation of a predictive model/algorithm for DILI assessment was out of our scope. According to publicly available information, Novartis is developing an algorithm called the ‘DILI cluster score’ using pharmacokinetic and physiochemical properties to predict the risk for orally administered drugs [37].

6 Conclusion

The over-reporting of severe life-threatening hepatic reactions, including autoimmune hepatitis, associated with UPA in post-marketing surveillance may be partially explained by its physiochemical (high lipophilicity) and pharmacokinetic (hepatic metabolism, long half-life, inhibition of liver transporters, reactive metabolite formation) features. These findings indicate the need for further investigations to clarify the mechanistic basis of DILI, which remains unsatisfactory, and underline the unpredictable nature of liver damage in clinical practice. We call on clinicians to raise awareness about the rare occurrence of DILI with UPA even in patients without apparent risk factors. This “bedside-to-bench” pharmacological approach may aid regulators in exploring the underlying mechanisms subtending DILI recognised in post-marketing surveillance, thus supporting issued regulatory warnings.

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Declarations

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Conflicts of interest MG, EP, FDP, and ER have no conflicts of interest that are directly relevant to the content of this article.

Ethics approval Not applicable.

Consent to participate Not applicable.

Consent for publication Not applicable.

Availability of Data and Material Data supporting the findings of this study were derived from the following resource, which is available in the public domain: https://fis.fda.gov/sense/app/d10be6bb-494c-4cd2-82e4-0135608ddc13/sheet/7a47a261-d58b-4203-a8aa-6d3021737452/state/analysis.

Code availability Not applicable.

Table 2 (continued)

| Liver Injury Associated with Ulipristal Acetate |
|-----------------------------------------------|
| **Liver Injury** | **Associated with Ulipristal Acetate** |
| Glutathione conjugate of mono-oxygenated ulipristal acetate. This proposed structure is consistent with the oxidation of the 4,5 carbon atoms to a reactive epoxide, followed by deactivation through glutathione conjugation |
| A score close to 1 indicates a high probability of causing DILI, hyperbilirubinemia, or cholestasis. A score close to 0 indicates a high probability of not causing DILI, hyperbilirubinemia, or cholestasis |
| Drugs receiving a score >7, between 3 and 7, and <3 show, respectively, most DILI concern/solid evidence, less DILI concern/weak evidence, and no DILI concern/no evidence of hepatotoxicity |

| **Cluster Score** | **Properties** |
|-------------------|----------------|
| Glutathione conjugate of mono-oxygenated ulipristal acetate. This proposed structure is consistent with the oxidation of the 4,5 carbon atoms to a reactive epoxide, followed by deactivation through glutathione conjugation |
| A score close to 1 indicates a high probability of causing DILI, hyperbilirubinemia, or cholestasis. A score close to 0 indicates a high probability of not causing DILI, hyperbilirubinemia, or cholestasis |
| Drugs receiving a score >7, between 3 and 7, and <3 show, respectively, most DILI concern/solid evidence, less DILI concern/weak evidence, and no DILI concern/no evidence of hepatotoxicity |
Author Contributions ER made substantial contributions to the conception and design of the study. MG made substantial contributions to the acquisition and analysis of data. MG, ER, EP, and FDP made substantial contributions to interpretation of the data. MG was involved in drafting the manuscript. ER, EP, and FDP made substantial contributions to critically revising the manuscript for important intellectual content. All authors approved the final version of the manuscript.

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