

How does ITP impact the patient and their HRQoL?

Fatigue

Venous thromboses

Heavy menstrual bleeding

Unmet medical need

- Fatigue affects HRQoL and has significant socioeconomic consequences¹
- Fatigue affects 22–45% of patients with ITP²
- Causes not fully understood¹

Unmet medical need

- Adults with ITP have greater risk of TE vs general population⁴
- Estimated VT incidence in ITP population:
 0.41–0.67 per 100 person-years⁵

Unmet medical need

- ITP may cause **HMB**, which can **impact QoL**⁶
- Estimated **prevalence of HMB** in patients with ITP is 6–55% at **diagnosis** and 17–79% **during disease**⁶
- HMB may cause iron deficiency or IDA⁶

Monitoring

- Check fatigue is not caused by:3
 - Low iron
 - · Thyroid function problems
 - Depression
- PRO tool: FACIT-F questionnaire1

Monitoring⁵

- Monitor platelet count
- Thrombotic risk factors include:
 - Older age
 - Secondary ITP
 - Multiple prior ITP therapies
 - Use of TPO-RAs

Monitoring⁶

- Clinical criteria can include: cycle of
 ≥7 days; changing protection at least every
 2 hrs or at night; clots; iron deficiency;
 impacting social participation
- PRO tools: **PBAC, ITP-PAQ, MMAS**

Management³

- Support from ITP patient groups
- Psychosocial support includes:
 - Regular exercise
 - Healthy eating
 - Reducing stress
 - · Balancing home-work-life
 - Talking to family/friends

Management

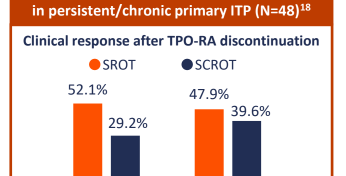
- No standard treatment guidelines⁵
- Treatments include:4
 - Antithrombotics e.g. warfarin, LMWH, DOAC
 - Anticoagulants + antiplatelet

Management

- Limited options that do not permanently impair fertility:
 - Antifibrinolytics ± hormonal therapy
- Options that permanently impair fertility:
 - Endometrial ablation; hysterectomy⁶
- Iron supplementation for iron deficiency/IDA⁷



What are the latest data that inform the use of current treatments for chronic ITP?



Long-term follow-up of the STOPAGO study

~50% of patients maintained SROT for >4 years after TPO-RA discontinuation

Long-term FU

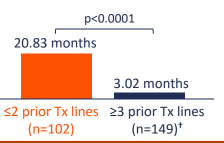
(>4 years)

Main study

(1 year)

Outcome of splenectomy after the year 2000 in patients with refractory ITP from the UK ITP Registry²⁰

Median time to first treatment post-splenectomy*



Probability of splenectomy inducing sustained remission in refractory patients with ITP is considerably lower vs non-refractory patients

First line

Corticosteroids⁸ IVIg / Anti-D Ig^{8,9}

Second line onwards

TPO-RAs

Eltrombopag^{8,9,10,11} Romiplostim^{8,9,12,13} Avatrombopag^{8,9,14,15}

Anti-CD20

Rituximab (off label)8,9

SYK inhibitor

Fostamatinib^{9,16,17}

Other immunosuppresants^{8,9}

Splenectomy^{8,9}

Italian real-world experience with fostamatinib in adult patients with refractory symptomatic chronic ITP¹⁹







Overall response within 3 months

Complete response[‡]

Still receiving Tx at 6 months

59 side effects reported in 38 patients (31 treatment related); most common were: diarrhoea (n=13), hypertension (n=8), transaminitis (n=8), neutropenia (n=4)

Data suggest fostamatinib is an effective therapeutic option in real-world practice with an acceptable safety profile

[†]Refractory ITP defined as having received ≥3 lines of therapy. [‡]According to International Working Group criteria.



^{*}Median time to first treatment post-splenectomy was a surrogate marker of splenectomy failure.

What data are there for emerging chronic ITP treatments?



~1 in 5 patients with ITP fail to achieve a platelet count >50 x 10⁹/L after treatment with available therapies or may encounter loss of response or intolerance. A significant disease burden remains with an unmet medical need to find a well-tolerated, targeted disease-modifying therapy²¹

Treatment and MoA



Rilzabrutinib
BTK inhibitor²³

Avatrombopag (paediatric use)
TPO-RA²⁵

Mezagitamab

CD38 inhibitor²¹

Phase

arms



Phase III

Randomized 2:1 sovleplenib (n=126) vs placebo (n=62) 300 mg QD

ESLIM-01²²

Key efficacy results

Treatment



DRR: 48% vs 0% (p<0.0001)*

ORR (all p<0.0001)

- ≥1 PC ≥50 x 10⁹/L: 71% vs 16%[†]
- Two consecutive PCs ≥30 x 10°/L and double from BL: 73% vs 6%
- PC ≥30 x 10⁹/L and increased ≥20 x 10⁹/L from BL: 75% vs 22%[‡]

Key safety results



TEAEs: 99% vs 85%

Grade 3/4 TEAEs: 25% vs 24%

Most common TEAEs: URTI, COVID-19, ↑ blood LDH

GI toxicities: Nausea 1.6% vs 3.2%; vomiting 1.6% vs 1.6%; diarrhoea 1.6% vs 0%

Thromboembolic events: 0%

LUNA 2^{23,24}

Phase II

Rilzabrutinib (N=71) 400 mg BID

Pooled outcomes²³

• Durable response: 28%§

• Overall response: 41%

• Complete response: 35%¶

Long-term outcomes²⁴

- n=8/17 discontinued ≥1 or ↓
 concomitant ITP therapy
- Visits reaching median PC of ≥50 x 10⁹/L: 90%

All AEs: 86%²³ (LT data: 81%)²⁴ **TRAEs:** 61%²³ (LT data: 41%)²⁴

Grade ≥3 AEs: 17% (all TRAEs

grade 1/2)²³

Most common TRAEs:

Diarrhoea; nausea; headache; fatigue; vomiting²³

Thromboembolic events: 0%23,24

AVA-PED-301²⁵

Phase III

Randomized 3:1 avatrombopag (n=54) vs placebo (n=21) 10 or 20 mg QD (age dependent)

- DPR: 27.8% vs 0% of patients (p=0.0077)**
- PR: 81.5% vs 0% of patients (p<0.0001)*[†]
- PC ≥50 x 10⁹/L: 48.9% vs 1.2% of weeks (p<0.0001)*[‡]
- PC ≥50 and ≤150 x 10⁹/L: 29.2% vs 1.2% of weeks (p<0.0001)*[‡]

TEAEs: 92.6% vs 76.2%

TRAEs: 13.0% vs 4.8%

Most common TEAEs:

Petechiae; epistaxis; bruising;

headache

Thromboembolic events: 0%

NCT04278924^{21,26}

Phase II

Randomized mezagitamab (n=28) vs placebo (n=13) 100, 300 or 600 mg QW

Mezagitamab 100/300/600 mg vs placebo

- PR: 66.7/62.5/90.9% vs 23.1%*§
- Complete PR: 55.6/50.0/81.8% vs 0%*||
- Clinically meaningful PR: 66.7/75.0/90.9% vs 30.8%*¶
- Haemostatic PR: 40.0/25.0/100% vs 0%***

TEAEs: 67.9% vs 69.2%

TRAEs: 32.1% vs 38.5%

Grade ≥3 TEAEs: 17.9% vs

23.1%

*PCs $\geq 50 \times 10^9/L$ at 4–6 visits during 14–24 weeks, not impacted by rescue treatment. $^{\dagger}Not$ impacted by rescue treatment. $^{\ddagger}For$ patients with a PC <15 x 10 $^9/L$ at BL. $^{\$}\geq 8$ of the last 12 PCs $\geq 50 \times 10^9/L$. $^{\$}\geq 2$ consecutive PCs $\geq 50 \times 10^9/L$ and increased $\geq 20 \times 10^9/L$ from BL. $^{\$}PCs \geq 100 \times 10^9/L$. **PC $\geq 50 \times 10^9/L$ in 12-week core phase in absence of rescue therapy. * $^{\$}\geq 2$ consecutive PC $\geq 50 \times 10^9/L$ in 12-week core phase in the absence of rescue therapy. * $^{\$}D$ visits. $^{\$}PC \geq 100 \times 10^9/L$ on ≥ 2 visits. * $^{\$}PC \geq 20 \times 10^9/L$ above BL on ≥ 2 visits. * $^{\$}PC \geq 100 \times 10^9/L$ above BL on ≥ 2 visits.



Abbreviations and references

Abbreviations

AE, adverse event; BID, twice daily; BL, baseline; BTK, bruton tyrosine kinase; CD, cluster of differentiation; DOAC, direct oral anticoagulant; DPR, durable PR; DRR, durable response rate; EHA, European Hematology Association; EMA, European Medicines Agency; FACIT-F, Functional Assessment of Chronic Illness Therapy-Fatigue; FDA, United States Food and Drug Administration; FU, follow-up; GI, gastrointestinal; HMB, heavy menstrual bleeding; HRQoL; health-related QoL; IDA, iron deficient anaemia; Ig, immunoglobulin; ISTH, International Society on Thrombosis and Haemostasis; ITP, immune thrombocytopenia; IVIg, intravenous Ig; LDH, lactate dehydrogenase; LMWH, low molecular weight heparin; LT, long term; MoA, mechanism of action; MMAS, menorrhagia multi-attribute scale; ORR, overall response rate; PAQ, Patient Assessment Questionnaire; PBAC, pictorial blood loss assessment chart; PC, platelet count; PR, platelet response; PRO, patient-reported outcome; QD, once daily; QoL, quality of life; QW, once weekly; SCROT, sustained complete response off treatment; SROT, sustained response off treatment; SYK, spleen tyrosine kinase; TE, thromboembolism; TEAE, treatment-emergent AE; TPO-RA, thrombopoietin receptor agonist; TRAE, treatment-related AE; Tx, treatment; URTI, upper respiratory tract infection; VT, venous thrombosis.

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