



Source: L	SEG, 2024
-----------	-----------

Market data	
EPIC/TKR	STX/SHIEF
Price (p)	4.9/0.01
12m high (p)	12.5/0.15
12m low (p)	1.1/0.015
Shares (m)	782.1
Mkt cap (£m)	37.9
EV (£m)	53.2
Free float*	59%
Country of listing	UK
Reporting currency	USD
Market	AIM
*As defined	by AIM Rule 26

Description

Shield is a specialty pharma company with an iron replacement product, Feraccru/ACCRUFeR, approved in the US and Europe for the treatment of iron deficiency in adults, with or without anaemia. In the US, ACCRUFeR is co-marketed by Shield and Viatris, with the aim of rapidly expanding its market share. In Europe, Feraccru is marketed by Norgine, with Shield receiving royalties.

Company inf	ormation
CEO	Anders Lundstrom
CFO	Santosh Shanbhag
Chairman	Hans Peter Hasler

+44 (0)191 511 8500 www.shieldtherapeutics.com

Key shareholders	
Directors AOP Health Hargreaves Lansdow Nestle SA Interactive Investor AJ Bell	1.5% 39.8% n 9.1% 7.2% 6.9% 4.5%
Diary	
Oct'24 Jan'25 Mch'25	3Q'24 Rx data Trading update 2023 results
Analyst	
Martin Hall mh@ł	ardmanandco.com

SHIELD THERAPEUTICS

Focusing on successful execution

Shield is a commercial-stage pharma company delivering specialty products that address the unmet medical need of patients with iron deficiency (ID). Since its July 2021 US launch, Shield and Viatris have increased physician awareness of the differentiating characteristics of ACCRUFeR[®] as an oral ID drug, in order to generate sales traction. 1H'24 results have confirmed that sales continue to progress well, while costs have been closely controlled. Management is focused on successful execution and reaffirmed its view that the gross cash position coupled with expected growth will be sufficient to see Shield through to cashflow-breakeven in 2H'25.

- Strategy: Shield and co-marketing partner Viatris are commercialising ACCRUFeR in the US. Elsewhere, Shield's strategy is to out-license commercial rights to partners with appropriate expertise in target markets, which has been achieved so far in Europe, China, Republic of Korea and Canada.
- Interims: In 1H'24, sales grew 224% to \$12.13m (\$3.74m) on the back of strong ACCRUFeR Rx (+160%), improved pricing (+33%) and better-than-expected performance by partners (EU +80%). Consequently, the underlying EBIT loss, at \$14.1m, was about \$1.0m better than forecast.
- Cash management: Shield had already reported, in a trading update, that the gross cash was better than previously forecast, at \$8.1m on 30 June. This excludes both the \$5.7m China milestone monetisation deal with major shareholder, AOP Health, and the £0.25m/\$0.32m Canada approval milestone.
- ▶ Forecasts: Recent statements have indicated a subtle change in emphasis towards ACCRUFeR sales and away from Rx growth (influenced by consignment volume) and Rx pricing (lower levels of subsidised Rx). Our full-year 2024 sales forecast has been reduced by a modest \$1.0m, to \$31.5m, to reflect this change.
- Investment summary: The interim results confirmed that Shield is continuing to move in the right direction and the new CFO has a firm grip on the cash position. ACCRUFeR Rx growth, coupled with improved pricing (lower discounting), is generating sales that are above previous (Jan'24) forecasts. While the beneficial 1H'24 working capital position is likely to unwind in 2H'24, receipt of recent milestones provides further flexibility to cash management.

Financial summary and valuation								
Year-end Dec (\$m)	2020	2021	2022	2023	2024E	2025E		
Product sales	0.94	1.40	5.50	13.09	31.50	73.12		
R&D	-3.31	-0.80	-1.32	-1.81	-1.30	-0.50		
Other income	11.40	0.79	1.11	4.41	6.05	1.90		
EBITDA	-0.29	-24.46	-28.55	-30.26	-18.34	-4.81		
Underlying EBIT	-3.77	-27.50	-31.40	-31.33	-19.54	-6.01		
Underlying PBT	-3.76	-27.50	-30.99	-31.83	-22.15	-7.82		
Underlying EPS (¢)	-3.20	-13.27	-13.49	-4.52	-2.83	-1.00		
Statutory EPS (¢)	-3.79	-13.11	-21.25	-4.59	-2.83	-1.00		
Gross cash	4.01	16.33	3.40	13.95	4.82	3.17		
Net cash/(debt)	3.97	16.12	-3.95	-6.30	-15.18	-21.83		
Equity issues	0.01	40.21	2.67	28.16	0.00	0.00		
EV/sales (x)	-	-	-	4.2	1.7	0.7		

Table of contents

1H'24 results	3
Operational highlights	3
Financial highlights	3
Operational update	5
Progress against 2024 business objectives	5
ACCRUFeR	5
Global partnerships	8
Financing update	9
Financials and investment case	
Income statement	
Balance sheet	
Cashflow	
Valuation	
Company matters	
Disclaimer	
Status of Hardman & Co's research under MiFID	



1H'24 results

Operational highlights

- ► ACCRUFeR: Shield reported a 160% increase in ACCRUFeR Rx in 1H'24. Given that 1Q'24 growth was broadly flat on 4Q'24, due to the well-documented change in the Texas PBM, this growth was driven by an excellent 2Q'24, reflecting the 12-month anniversary of the expanded and trained US sales team. Continued sequential quarterly growth is expected to continue moving forward.
- ► ACCRUFeR discounts: There is clear evidence that the efforts being made by the Shield-Viatris sales team to reduce the level of subsidised Rx are being rewarded, with the average Rx generating \$139 in 1Q'24, and \$171 in 2Q'24. Further progress is expected as the commercial team continues to lower the level of consignment volumes.
- Partnering: Progress has been made during 1H'24 with several partners. The sales performance of Norgine in Europe was better than expected, Kye received ACCRUFeR regulatory approval in Canada, Korea filed for regulatory approval in Korea, and ASK continued to enrol patients into its Phase III regulatory trial.

Financial highlights

- Sales: ACCRUFeR generated sales of \$10.96m (+148%) through a combination of Rx growth and lower discounts. This was supported by European royalties of \$1.07m (+80%) and modest sales from Korea. Total sales were \$12.13m (+224%).
- ► **COGS:** Manufacturing and supply costs were below expectations, generating a gross margin of 45.0% (44.3%). While similar to 1H'23, it was considerably higher than the gross margin for the whole of 2023. The COGS is largely dictated by the percentage of US sales payable to Viatris and the 5% royalty.
- SG&A: Despite the cost of employing the enlarged sales team for the whole of 1H'24, the underlying SG&A costs increased only 8% to \$18.23m (\$16.89m). Underlying SG&A is expected to be around this level again in 2H'24.
- ▶ Net cash/(debt): At 30 June, Shield had gross cash of \$8.1m and net debt of \$11.87m. Financial flexibility has been enhanced by a receivables financing deal with Sallyport of up to \$10m, of which \$6.4m was drawn down on 30 June. Flexibility has been further enhanced through the China milestone monetisation agreement with AOP (+\$5.7m) and regulatory approval in Canada (+\$0.32m).

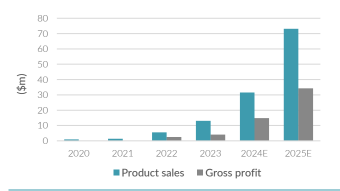
Interim results summary – actual vs. expectations								
Year-end Dec	1H'23	1H'24	Growth	1H'24	Delta			
_(\$m)	actual	actual	CER	*forecast	Δ			
Product sales	4.33	12.13	+224%	11.53	+0.60			
COGS	-2.09	-6.68	n/m	-6.92	+0.24			
SG&A	-16.89	-18.23	+8.0%	-18.55	+0.32			
Share-based costs	-0.18	-0.58	n/m	-0.44	-0.14			
R&D	-0.43	-0.75	+73%	-0.65	-0.10			
Other income	4.30	0.00	-	0.00	-			
Underlying EBIT	-11.54	-14.06	+22%	-15.03	+0.97			
Gross cash	13.59	8.10	-	6.85	+1.25			
Debt/leases	-5.77	-19.97	-	-20.00	+0.03			
Net cash/(debt)	7.82	-11.87	-	-13.15	+1.28			

*Prior to release of business update statement on 24 July Note: numbers may not add up exactly due to rounding Source: Hardman & Co Research

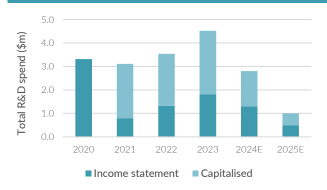
Shield Therapeutics



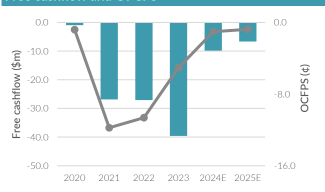
Product sales and gross profit



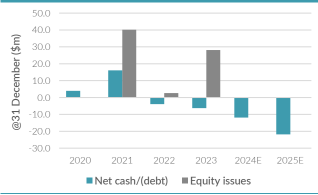
Total R&D investment



Free cashflow and OFCPS



Net cash/(debt) and equity issues



- ► The US is the only market in which Shield records product sales. In other countries, it partners with distributors.
- Shield is recording 100% of net sales of ACCRUFeR in the US, with Viatris's 45% proportion being charged to COGS, reducing the gross margin from 85% to ca.45%.
- Sales milestones and royalties from commercial partners are treated as "other income", and are not included in our sales.
- Total R&D spend in 2023 peaked at \$4.5m. However, only 40% was recorded through the income statement; the other 55% was capitalised.
- Main investment is a paediatric study with Feraccru/ ACCRUFeR initiated at the end of 2021; this was forecast to cost \$6.5m-\$7.0m over a three-year period. Most of these costs are now behind the group, allowing R&D spend to ease off.
- Shield has been cashflow-negative during the investment phase and commercialisation of Feraccru/ACCRUFeR.
- 2024 cashflows will continue to be dominated by ramp-up in ACCRUFeR marketing spend for the co-promotional programme agreed with Viatris.
- Rapid acceleration of ACCRUFeR Rx and sales in 2024 and 2025 are expected to see Shield become cashflowbreakeven in 2H'25 and beyond.
- Gross cash at 30 June 2024 was \$8.1m, offsetting \$20.0m of debt.
- More favourable covenants on the SWK debt facility were negotiated in April 2024.
- Shield arranged a flexible accounts receivable facility up to \$10m with Sallyport in April to provide greater working capital flexibility.
- ▶ In July 2024, Shield monetised its potential China regulatory milestone to raise an additional \$5.7m to strengthen the balance sheet further.

Source: Company data; Hardman & Co Research

Progress against three clearly defined goals

Operational update

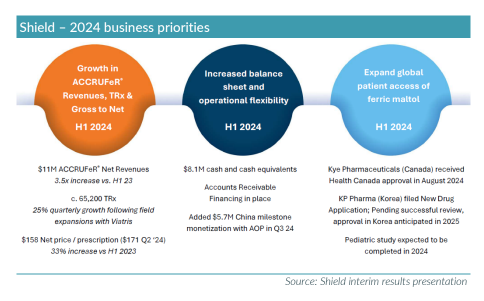
Progress against 2024 business objectives

HARDMAN&CO.

In its presentation of fiscal 2023 results, management set out three clear business objectives for 2024:

- Maximise revenues (product sales and royalties) directly in the US through the commercial deal with Viatris and indirectly through its overseas partners. In the US, the aim was to grow ACCRUFeR RX and reduce the discounting and subsidies frequently used in the industry when a drug is first used.
- Increase balance sheet and operational flexibility through tight control of operating expenses and enhancements to working capital. A clear focus on making investments that were directly tied to supporting ACCRUFeR.
- Expand global patient access to ferric maltol by working closely with overseas partners.

The following slide, extracted from Shield's interim results presentation, shows that the company has made considerable progress against all three business objectives in 1H'24, which has continued further so far in 2H'24.



ACCRUFeR

ACCRUFeR Rx progress

2024 did not start well for Shield when the largest state (Texas) user of ACCRUFeR decided to change its Pharmacy Benefit Manager (PBM), throwing the whole process of pre-approvals into disarray for a short period until a new PBM was appointed from April. Really good performances in other states – notably New York and California – only just offset the impact from Texas, with 1Q'24 ACCRUFeR Rx rising 1%.

2Q'24 saw the Texas State Medicaid programme return to normal and, with continued growth elsewhere, ACCRUFeR Rx grew 143% compared with 2Q'23 and 26% over 1Q'24. The top states for ACCRUFeR are currently California, Florida, New York, and Texas.

Good recovery in 2Q'24 after poor start in 1Q'24 in Texas

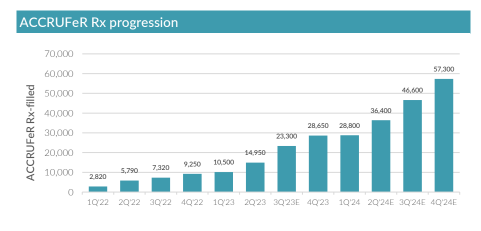
Return to sequential quarterly Rx growth



Shield Therapeutics

Continually addressing the volume:price balance

One characteristic of the large state Medicaid programmes is something called consignment volume, whereby higher discounts are demanded for high volumes. During 2Q'24, Shield-Viatris has continued its goal to redress the balance between price and volume, gradually reducing the number of subsidised Rx.

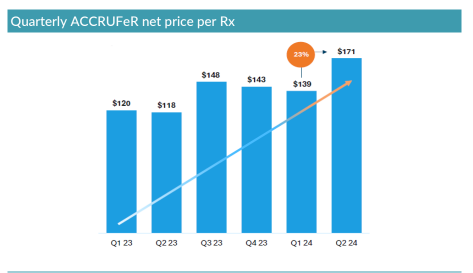


Source: Hardman & Co Research

ACCRUFeR pricing trend

In order to increase awareness of ACCRUFeR among physicians and to establish the drug in the market, ACCRUFeR was given away free, or heavily subsidised, when it was first launched. This is normal industry practice. As new products become better known, more established, and included under Medicaid and Medicare formulary coverage, the aim is to reduce the discount levels. A key goal of the Shield-Viatris commercial team over the past 12 months has been to reduce the average discount that each ACCRUFeR Rx attracted.

During 1H'24, and particularly in 2Q'24, Shield has made significant progress in reducing these discounts with the long-term aim of achieving an average price per Rx of \$220. The following chart shows the average net ACCRUFeR price, on a quarterly basis, since 1Q'23. In the same way that Rx levels in the industry tend to be lower in the early part of each year, pricing also tends to be lower. This can be seen in the dip between 4Q'23 and 1Q'24. However, in 2Q'24, Shield recorded its best quarter for pricing with the average price reaching \$171. Further progress on this goal is expected in 2H'24.



Source: Shield interim results presentation

2Q'24 Rx price outcome of \$171 was

best to date

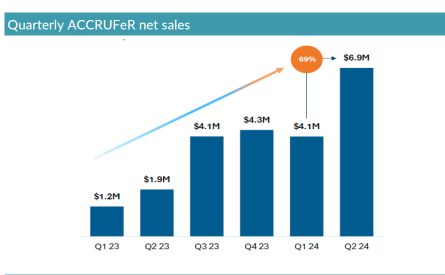


Sales beginning to dominate the focus

ACCRUFeR sales

While ACCRUFeR Rx growth and average net prices remain very important and are both contributory factors, a general observation that we have made in recent trading updates and results announcements is a subtle change in emphasis on to sales.

Clearly, sales are dependent on both Rx levels and price, but, moving forward, we envisage that Shield-Viatris is trying to move away from notching up an Rx at any price, thereby improving the quality of each Rx filled. This might result in a slight reduction in the quarterly Rx growth rate reported but at continually improving average price. Consequently, we see greater focus on sales and sales growth. To that end, the company provided accurate back information on quarterly ACCRUFeR sales, corrected for all the well-documented issues in 2023.



Source: Shield interim results presentation

In our opinion, this is also a reflection of the desire of the new CFO to put his stamp on the quality and consistency of the information being provided in recent trading updates and financial reports. There is no doubt that the 1H'24 report and presentation had much greater transparency and that the data can be relied upon by investors.

Changes to ACCRUFeR forecasts

On the basis that Shield-Viatris is trying to reduce the level of consignment volume and improve process, we have tweaked downwards our growth expectations for ACCRUFeR Rx but slightly raised our average price per Rx. Although this has resulted is a slight fall in US ACCRUFeR sales, it is compensated at the group level by increased forecast for the contribution from Norgine in Europe.

Changes to ACCRUFeR forecasts									
	1Q'24	2Q'24	3Q'24E	4Q'24E	2024E				
ACCRUFeR Rx growth rate									
Old	+1%	+26%	+31%	+30%	174,800				
New			+28%	+23%	169,100				
ACCRUFeR average net Rx									
price									
Old	\$130	\$171	\$175	\$179	\$170				
New			\$176	\$181	\$173				
ACCRUFeR net sales									
Old	\$4.1m	\$6.9m	\$8.4m	\$11.1m	\$30.5				
New			\$8.1m	\$10.2m	\$29.3				
			Source	Hardman &	Co Research				

Data quality has improved under new CFO

Slight tweak to forecasts due to volume versus price balance



Global partnerships

To date in 2024, Shield has made progress with each of its non-US partners, which are summarised in the following graphic.



Source: Shield interim results presentation

Norgine

The slow progress of Norgine – EU, UK, Norway, Australia, New Zealand plus some other non-EU Countries – has been a frustration, particularly in Europe. However, progress does appear to be evolving, as evidenced by the 80% increase in royalties received in 1H'24, which was much greater than forecast. After some initial problems with NICE and the payors in the UK, Feraccru is now being accepted on many more hospital formularies. We expect a continuation of this progress in 2H'24.

Kye

Kye filed for approval of ACCRUFeR with the Canadian regulator in 2023 and approval was always anticipated in 2H'24. This duly arrived at the end of August, triggering an approval milestone of ± 250 k/\$325k, which is due to be paid imminently. Kye is now going through the pre-launch phase and is expected to formally launch in Canada in 1Q'25.

Korea Pharma

Following the successful completion of the required pharmacokinetic study, Korea Pharma has submitted ACCRUFeR to the regulator for approval in the Republic of Korea. A decision is expected during 2025.

ASK Pharma

For approval in China, ASK is required to undertake a pivotal Phase III clinical trial. The protocol is similar to those used to obtain regulatory approval from the EMA and the FDA. Although, as discussed previously, recruitment has been slower than anticipated, good progress has been made in 1H'24 and full enrolment is targeted for completion towards the end of 2024. Trial results are anticipated in 2025 and, if successful, the regulatory submission is also expected to occur next year.

Paediatric study

As part of the regulatory process in Europe and the US, Shield was required to undertake a Phase III paediatric study (FORTIS/ST10-01-305) comparing the safety, tolerability and effectiveness of an oral liquid suspension of ferric maltol with oral ferrous sulphate liquid in children with iron deficiency anaemia. This trial is expected to complete by the end of 2024.

Imminent receipt of £0.24m/\$0.32m milestone



Financing update

Since his appointment at the end of 2023, the new CFO has undertaken a complete review of the company's finances and spending plans. As stated earlier, in our opinion, there has been a noticeable improvement in the quality of the information being provided and that investors can rely on these data.

He inherited a stretched balance sheet, with the market generally of the view that more capital was required. However, through a series of operational decisions and changes and some financing deals, the balance sheet has improved and the current and forecast gross cash position is expected to see the company through to cashflow-breakeven in 2025, assuming that internal and external forecasts are met.

Operating costs

The first thing that the new CFO addressed was to reschedule some of company's operating costs to ensure that investment was in a focused and timely manner to grow ACCRUFeR in the US.

In 1H'24, US selling costs rose 10.2%, reflecting the full contribution from the expanded sales team, which was only in the recruitment phase in 1Q'23. General administration costs (excluding share-based payments) were reduced by 1.4% to \$5.12m (\$5.19m), despite the negative impact of translating the UK administrative costs into USD. D&A rose substantially in 1H'24. Taking all of these points into account, the underlying SG&A costs rose only 8.0% in 1H'24. Management maintained, at the analyst meeting, that costs would be largely similar in 2H'24 compared with 1H'24, but we have prudently allowed for ca.\$1.5m increase in our forecasts.

SG&A costs						
\$m	1H'23	2H'23	2023	1H'24	2H'24E	2024E
US selling costs	-11.20	-10.52	-21.72	-12.34	-13.38	-25.72
General administration	-5.19	-9.11	-14.30	-5.12	-5.55	-10.67
D&A	-0.50	-0.58	-1.07	-0.77	-0.77	-1.54
Underlying SG&A	-16.89	-20.20	-37.09	-18.23	-19.69	-37.93
Share-based costs	-0.18	-0.70	-0.88	-0.58	-0.58	-1.17
Reported SG&A	-17.06	-20.90	-37.96	-18.82	-20.28	-39.10

Source: Hardman & Co Research

It should be noted that the general administration costs were abnormally high in 2H'23. This was due to the inclusion of various financing costs within this figure, which are not expected to be repeated.

Financing deals

Shield has undertaken three financing/refinancing deals so far in 2024 to strengthen the balance sheet and provide increased flexibility to its working capital.

SWK Funding LLC

Shield renegotiated its existing \$20m debt financing agreement with SWK to provide more favourable covenant terms. On a rolling 12-month basis, the minimum ACCRUFeR sales targets are \$16.5m, \$22.5m, \$31.5m, \$38.9m, and \$45.7m in 2Q'24, 3Q'24, 4Q'24, 1Q'25, and 2Q'25 and beyond, respectively. Shield maintained that its internal forecasts would consistently comply with these revised minimum sales requirements.

Hands-on control of operating cost

Revised sales covenants are on a rolling 12-month basis





Source: Shield interim results presentation

Sallyport Commercial Finance

In April 2024, Shield strengthened its balance sheet through an accounts receivable arrangement with Sallyport for up to \$10m. On issue of an invoice in the US, Shield can draw down the receivable from Sallyport, which is then repaid, less costs, to Sallyport, immediately upon payment receipt from the customer. This improves Shield's working capital position, covering, for example, the average 72-day payments from its largest customer, Cardinal Health.

The amount drawn down will vary on a day-to-day basis. On 30 June, Shield had an outstanding draw down of \$6.84m with Sallyport, showing up in short-term liabilities (note 11 in the accounts). In addition, under the terms of the agreement, Shield is required to hold \$1.0m of restricted cash as a contingency until closure of the agreement, listed under long-term assets.

AOP Health

On 3 July, Shield announced that it had signed an agreement with its major shareholder, to monetise the anticipated \$11.4m China approval milestone from ASK Pharma. Under the terms of this agreement, AOP has paid Shield \$5.7m immediately in return for the right to receive the full \$11.4m from ASK within 30 days from the "approval milestone" being achieved. ASK is expected to complete enrolment into the Phase III trial by the end of 2024 and, all being well, the approval milestone is expected to become payable around the end of 2026.

In the event that the approval milestone has not been triggered by 31 December 2026, or in the event that this agreement is terminated, including at Shield's election or due to a breach by Shield of its terms, the \$5.7m advance plus accrued interest and fees at an interest rate of SOFR+9.25% (calculated from the date of the advance until the day of payment) and an exit fee of 6.5% of the advance will be payable by Shield to AOP.

Summary

Management is totally focused on commercial execution. All of these financing deals, together with close control of costs and working capital are designed to see Shield through to cashflow-breakeven in 2H'25.

Accounts receivable facility will vary on a day-to-day basis...but up to \$10m

Accounting for AOP arrangement still uncertain...but we have included whole \$5.7m in "other income"



Financials and investment case

Income statement

- Sales: Strong performance in 1H'24 and the key focus. Modest adjustment to 2024 and 2025 forecasts to reflect the balance between consignment volume and price with the aim of continually reducing the level of subsidised Rxs.
- COGS: There are three components to Shields's COGS apart from the basic product manufacturing costs, there are royalties and Viatris's 45% proportion of net sales. A normalised gross margin is around 45% of net sales.
- **SG&A:** Management intends to continue its planned investment into marketing in order to drive sales growth. However, it does have some flexibility around the timing of this investment to help with the management of its cash position.
- **R&D:** Shield is continuing to invest in the paediatric study required by the regulators. However, much of this R&D spend is capitalised.
- Other income: This is a combination of sales milestones from Viatris and regulatory milestones from global partners. 2024 includes \$5.7m received from AOP and the imminent approval milestone from Kye.
- Profitability: Shield is expected to become EBITDA-positive during 2H'25. Timing of profitability is dependent on receipt of sales milestones from Viatris, which look set to commence in fiscal 2026.

Income statement						
Year-end Dec (\$m)	2020	2021	2022	2023	2024E	2025E
Product sales	0.94	1.40	5.50	13.09	31.50	73.12
COGS	-1.74	-1.35	-3.04	-9.06	-16.70	-38.84
Gross profit	-0.80	0.05	2.46	4.03	14.80	34.27
Gross margin	7.2%	60.2%	44.7%	30.8%	47.0%	46.9%
SG&A (underlying)	-10.06	-26.19	-32.73	-37.09	-37.93	-40.52
Share-based costs	-0.99	-1.36	-0.91	-0.88	-1.17	-1.17
R&D	-3.31	-0.80	-1.32	-1.81	-1.30	-0.50
Other income	11.40	0.79	1.11	4.41	6.05	1.90
EBITDA	-0.29	-24.46	-28.55	-30.26	-18.34	-4.81
Depreciation	-0.03	-0.03	-0.04	-0.04	-0.04	-0.04
Amortisation	-3.45	-3.01	-2.81	-1.04	-1.17	-1.17
Underlying EBIT	-3.77	-27.50	-31.40	-31.33	-19.54	-6.01
Exceptional items	0.00	0.00	-18.11	0.00	0.00	0.00
Statutory EBIT	-3.77	-27.50	-49.51	-31.33	-19.54	-6.01
Net interest	0.00	0.01	0.41	-0.50	-2.61	-1.81
Forex gain/loss	0.27	0.38	0.00	-0.54	0.00	0.00
Underlying PBT	-3.76	-27.50	-30.99	-31.83	-22.15	-7.82
Extraordinary items	0.00	0.00	0.00	0.00	0.00	0.00
Statutory PBT	-3.50	-27.11	-49.10	-32.38	-22.15	-7.82
Tax payable/credit	-0.96	0.32	-0.46	-0.92	0.00	0.00
Underlying net income	-3.75	-27.13	-31.45	-32.75	-22.15	-7.82
Statutory net income	-4.45	-26.80	-49.56	-33.29	-22.15	-7.82
Ordinary 1.5p shares:						
Period-end (m)	117.62	215.89	259.39	782.06	782.06	782.06
Weighted average shares (m)	117.40	204.41	233.19	725.22	782.06	782.06
Fully diluted (m)	121.35	211.87	285.79	782.54	844.37	849.37
Underlying basic EPS (¢)	-3.20	-13.27	-13.49	-4.52	-2.83	-1.00
Statutory basic EPS (¢)	-3.79	-13.11	-21.25	-4.59	-2.83	-1.00
Underlying fully dil. EPS (ϕ)	-3.09	-12.80	-11.00	-4.19	-2.62	-0.92
Statutory fully dil. EPS (¢)	-3.67	-12.65	-17.34	-4.25	-2.62	-0.92
DPS (¢)	0.0	0.0	0.0	0.0	0.0	0.0
					rdman & Co	



Balance sheet

- ▶ Net cash/(debt): On 30 June 2024, Shield had gross cash of \$8.1m including the part drawdown of the accounts receivable financing deal. The \$20m debt facility from SWK was fully utilised. Post the period-end, Shield received the \$5.7m China milestone monetisation advance from AOP and payment of the Canada approval milestone is imminent.
- ▶ Loan facility: In September 2023, Shield entered into a \$20m five-year term loan facility with SWK Holdings. This term loan is interest-only (minimum 14.25%) for eight quarters, thereafter it will be interest plus \$1m capital repayment per quarter. Sales covenants were renegotiated to more favourable terms in April 2024.
- Inventories: Inventories increased 26% to \$4.0m in 1H'24 compared with the level at 31 December 2023. Shield has the flexibility to release some working capital at a suitable time point, again as part of the overall management of gross cash.
- ▶ Working capital: Overall, there was a \$3.90m release of working capital in 1H'24, but this includes the drawdown accounts receivable facility from Sallyport.

Balance sheet						
@31 Dec (\$m)	2020	2021	2022	2023	2024E	2025E
Shareholders' funds	41.33	55.31	6.54	15.03	-7.11	-14.93
Cumulated goodwill	0.00	0.00	0.00	0.00	0.00	0.00
Total equity	41.33	55.31	6.54	15.03	-7.11	-14.93
Share capital	2.41	4.36	5.37	15.01	15.25	15.25
Reserves	38.92	50.95	1.17	0.02	-22.36	-30.18
Provisions/liabilities	0.00	0.00	0.00	0.00	8.00	8.00
Deferred tax	0.00	0.00	0.00	0.00	0.00	0.00
Long-term leases	0.00	0.00	0.00	0.20	0.00	0.00
Short-term leases	0.04	0.21	0.11	0.21	0.00	0.00
Long-term loans	0.00	0.00	7.25	19.84	20.00	25.00
Short-term debt	0.00	0.00	0.00	0.00	0.00	0.00
<i>less</i> : Cash	4.01	16.33	3.40	13.95	4.82	3.17
less: Deposits	0.00	0.00	0.00	0.00	0.00	0.00
less: Non-core investments	0.00	0.00	0.00	0.00	0.00	0.00
Invested capital	37.35	39.19	10.49	21.33	16.07	14.90
Fixed assets	0.04	0.41	0.24	0.67	0.52	0.68
Intangible assets	37.22	36.20	14.21	16.86	17.35	16.87
Inventories	1.88	2.20	1.76	3.20	3.52	4.23
Trade debtors	0.30	1.10	4.09	9.99	10.93	12.08
Other debtors	0.55	2.85	2.40	3.51	3.51	3.51
Tax liability/credit	0.40	0.78	0.53	0.61	0.59	0.00
Trade creditors	-0.54	-1.77	-2.20	-4.05	-6.01	-6.99
Other creditors	-2.50	-2.58	-10.52	-9.47	-14.35	-15.47
Debtors less creditors	-1.79	0.38	-5.71	0.59	-5.33	-6.87
Invested capital	37.35	39.19	10.50	21.33	16.07	14.90
Net cash/(debt)	3.97	16.13	-3.95	-6.30	-15.18	-21.83
					rdman & Co	



Cashflow

- ▶ Net cash/(debt): Our forecasts assume that Shield will continue to fully utilise its \$20m SWK term loan facility. It will also continue to utilise its accounts payable facility, but this changes on a day-to-day basis based on the timing of the issuance of invoices and the payment receipt for those invoices. Careful management of the timing of planned marketing spend and working capital are expected to maintain a modest gross cash position at the end of each month and at the year end, obviating the need for an equity injection.
- ► Capitalised R&D: Although the paediatric study is continuing, much of the cost is contracted and pre-paid to the contract research organisation conducting the trial. A further \$0.98m was capitalised in 1H'24, but we expect this to tail off as the trial proceeds to a conclusion at the end of 2024.
- ▶ Working capital: Timing differences between supply of drugs to US wholesalers and payments have resulted in an increase in working capital, which has been offset by utilisation of accounts payable agreement with Sallyport. As ACCRUFeR Rxs accelerate and net selling discounts reduce, Shield will have more flexibility with its working capital to manage the gross cash position.

Cashflow						
Year-end Dec (\$m)	2020	2021	2022	2023	2024E	2025E
Underlying EBIT	-3.77	-27.50	-31.40	-31.33	-19.54	-6.01
Depreciation	0.03	0.03	0.04	0.04	0.04	0.04
Amortisation	3.45	3.01	2.81	1.04	1.17	1.17
Share-based costs	0.99	1.36	0.91	0.88	1.17	1.17
Inventories	1.20	-0.35	0.22	-1.45	-0.32	-0.70
Receivables	-0.34	-3.96	-2.79	-7.01	-0.94	-1.15
Payables	-2.66	2.26	7.27	1.91	9.96	0.98
Change in working capital	-1.80	-2.05	4.70	-6.55	8.70	-0.88
Exceptionals/provisions	0.00	0.00	0.00	0.00	3.40	0.00
Disposals	0.00	0.00	0.00	0.00	0.00	0.00
Other	0.34	0.71	-0.81	0.17	0.00	0.00
Company op. cashflow	-0.77	-24.44	-23.75	-35.76	-5.07	-4.52
Net interest	0.00	-0.04	-0.37	-0.10	-4.03	-1.81
Finance leases	-0.07	-0.11	-0.23	-0.09	-0.23	-0.23
Tax paid/received	-0.11	0.56	-0.43	-0.72	1.19	0.59
Operational cashflow	-0.94	-24.03	-24.78	-36.66	-8.14	-5.97
Capital expenditure	0.00	-0.51	-0.06	-0.24	-0.15	-0.19
Capitalised R&D	0.00	-2.32	-2.22	-2.71	-1.50	-0.50
Sale of fixed assets	0.00	0.00	0.00	0.00	0.00	0.00
Free cashflow	-0.94	-26.86	-27.06	-39.61	-9.79	-6.65
Dividends	0.00	0.00	0.00	0.00	0.00	0.00
Acquisitions	-0.03	-0.01	0.00	0.00	0.00	0.00
Disposals	0.00	0.00	0.00	0.00	0.00	0.00
Other investments	0.00	0.00	0.00	0.00	0.00	0.00
Cashflow after invests.	-0.97	-26.87	-27.06	-39.61	-9.79	-6.65
Share repurchases	0.00	0.00	0.00	0.00	0.00	0.00
Equity issues	0.01	40.21	2.67	28.16	0.00	0.00
Cost of fundraise	0.00	-2.09	-0.19	-1.35	0.00	0.00
Currency effect	0.82	0.53	-2.83	0.45	0.50	0.00
Loans/cash acquired	0.00	0.37	7.34	10.00	0.00	0.00
Change in net debt	-0.15	12.15	-20.07	-2.35	-9.29	-6.65
OCFPS (p)	-0.80	-11.76	-10.63	-5.06	-1.04	-0.76
Opening net cash/(debt)	4.12	3.97	16.12	-3.95	-6.30	-15.18
Closing net cash/(debt)	3.97	16.13	-3.95	-6.30	-15.18	-21.83
				Source: Ha	rdman & Cc	Research

Taking all recent events into account, valuation drops from 17p to 16p per share

Each quarterly update expected to see a continual rerating of shares

Covis acquired AMAG for \$647m in November 2020, and CSL bid CHF11.1bn for Vifor in December 2021 Valuation

Sum-of-the-parts

Our analysis, which pulls together the DCF models for each of the key commercial markets for Feraccru/ACCRUFeR, has been updated following all the recent events. This has generated a revised sum-of-the-parts EV of \$193m/£143m. The US valuation has reduced modestly along with the slightly reduced sales expectations. However, this has been more than offset by an increase in the European valuation following the better-than-expected performance from Norgine. The recent strength in sterling has negated the overall positive USD outcome. Updating the net debt position leaves our overall NPV per share at 16p, a 1p reduction.

Sum-of-the-parts valuation			
	NPV (I.c.)	NPV (GBP)	NPV per share
NPV of ACCRUFeR in US	\$117m	£90m	11p
NPV of Feraccru royalty stream in Europe	€33m	£28m	4р
Risk-adjusted NPV of China royalty stream	\$35m	£26m	Зр
Enterprise value	\$193m	£143m	18p
Net cash/(debt)	-\$21m	-£15m	-2p
Group valuation	\$173m	£128m	16p
Source: Hardman & Co Res			

Source: Hardman & Co Research

HARDMAN&CO.

Although there has been some recovery in the share price to ca.5.0p recently, the market is still factoring in the continuing commercial risk and the debate about gross cash, despite management maintaining its clear opinion that no equity issue is needed based on current projections. 1H'24 results showed that the PBM issue in 1Q'24 has gone away and there is real confidence in the numbers being presented. Further benefits of the enlarged sales team in each sequential quarter should give the market more confidence to continue to rerate the shares.

Comparator valuations

There are no directly comparable UK peers – so we continue to observe some global competitors for completeness. In addition, we have been highlighting the need of the pharma majors for M&A activity to bolster R&D pipelines and commercial portfolios. Iron replacement therapy is no exception. Covis (private company) paid an EV of \$647m in November 2020 for AMAG (EV/sales: 2.4x), and CSL bought Vifor for CHF11.1bn in December 2021 (EV/sales: 9.3x). In the event that ACCRUFeR targets are met in the next three years, the stance of Viatris would become very interesting, in our opinion, especially if the Shield valuation is slow to recover.

Comparative valuat	ion			
Company	*Vifor Pharma	*AMAG (now Covis)	Akebia Therapeutics	Shield Therapeutics
Ticker	VIFN	AMAG	AKBA	STX
Local currency	CHF	\$	\$	£
Share price	173.0	13.8	1.39	0.052
Shares in issue (m)	65.0	36.7	209.6	782.1
Market cap \$m)	11,245.0	505.0	290.3	69.8
Mkt cap (£m)	8,875.3	385.5	221.6	40.7
Cash	994.5	168.9	42.0	9.9
Debt	-603.6	-311.2	-37.8	-19.8
EV (\$m)	10,854.1	647.3	286.1	77.3
EV (£m)	8,566.8	494.1	218.4	59.0
EV relative to Shield	145.1	8.4	3.7	-

Share prices and currencies taken at close of business on 6 September 2024 *Based on shares in issue at date of completion of acquisition Source: Hardman & Co Life Sciences Research



Company matters

Registration

Incorporated in the UK with company registration number 09761509.

Registered office:

Northern Design Centre Baltic Business Quarter Gateshead Quays Newcastle NE8 3DF

+44 (0)191 511 8500

www.shieldtherapeutics.com

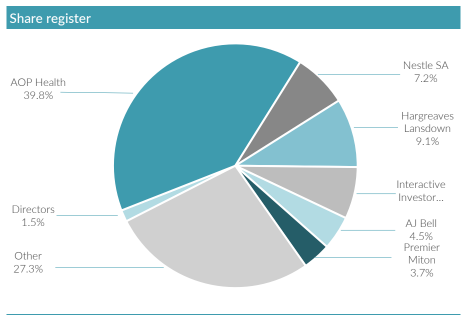
Board of Directors

Board of Directors				
Position	Name	Nominations	Remuneration	Audit
Chairman	Hans Peter Hasler	С	М	М
Chief Executive Officer	Anders Lundstrom			
Non-Executive Director	Fabiana Lacerca-Allen			М
Non-Executive Director	Peter Llewellyn-Davies	М		С
Non-Executive Director	Christian Schweiger	М		
Chief Financial Officer*	Santosh Shanbhag			

M = member; C = chair *Non-board PDMR appointment Source: Company reports

Share capital

On 6 September 2024, there were 782,056,367 Ordinary shares in issue. In addition, there are 57.32m options outstanding.



Shield Therapeutics



Disclaimer

Hardman & Co provides professional independent research services and all information used in the publication of this report has been compiled from publicly available sources that are believed to be reliable. However, no guarantee, warranty or representation, express or implied, can be given by Hardman & Co as to the accuracy, adequacy or completeness of the information contained in this research and they are not responsible for any errors or omissions or results obtained from use of such information. Neither Hardman & Co, nor any affiliates, officers, directors or employees accept any liability or responsibility in respect of the information which is subject to change without notice and may only be correct at the stated date of their issue, except in the case of gross negligence, fraud or wilful misconduct. In no event will Hardman & Co, its affiliates or any such parties be liable to you for any direct, special, indirect, consequential, incidental damages or any other damages of any kind even if Hardman & Co has been advised of the possibility thereof.

This research has been prepared purely for information purposes, and nothing in this report should be construed as an offer, or the solicitation of an offer, to buy or sell any security, product, service or investment. The research reflects the objective views of the analyst(s) named on the front page and does not constitute investment advice. However, the companies or legal entities covered in this research may pay us a fixed fee in order for this research to be made available. A full list of companies or legal entities that have paid us for coverage within the past 12 months can be viewed at http://www.hardmanandco.com/legals/research-disclosures. Hardman may provide other investment banking services to the companies or legal entities mentioned in this report.

Hardman & Co has a personal dealing policy which restricts staff and consultants' dealing in shares, bonds or other related instruments of companies or legal entities which pay Hardman & Co for any services, including research. No Hardman & Co staff, consultants or officers are employed or engaged by the companies or legal entities covered by this document in any capacity other than through Hardman & Co.

Hardman & Co does not buy or sell shares, either for their own account or for other parties and neither do they undertake investment business. We may provide investment banking services to corporate clients. Hardman & Co does not make recommendations. Accordingly, they do not publish records of their past recommendations. Where a Fair Value price is given in a research note, such as a DCF or peer comparison, this is the theoretical result of a study of a range of possible outcomes, and not a forecast of a likely share price. Hardman & Co may publish further notes on these securities, companies and legal entities but has no scheduled commitment and may cease to follow these securities, companies and legal entities without notice.

The information provided in this document is not intended for distribution to, or use by, any person or entity in any jurisdiction or country where such distribution or use would be contrary to law or regulation or which would subject Hardman & Co or its affiliates to any registration requirement within such jurisdiction or country.

Some or all alternative investments may not be suitable for certain investors. Investments in small and mid-cap corporations and foreign entities are speculative and involve a high degree of risk. An investor could lose all or a substantial amount of his or her investment. Investments may be leveraged and performance may be volatile; they may have high fees and expenses that reduce returns. Securities or legal entities mentioned in this document may not be suitable or appropriate for all investors. Where this document refers to a particular tax treatment, the tax treatment will depend on each investor's particular circumstances and may be subject to future change. Each investor's particular needs, investment objectives and financial situation were not taken into account in the preparation of this document and the material contained herein. Each investor must make his or her own independent decisions and obtain their own independent advice regarding any information, projects, securities, tax treatment or financial instruments mentioned herein. The fact that Hardman & Co has made available through this document various information constitutes neither a recommendation to enter into a particular transaction nor a representation that any financial instrument is suitable or appropriate for you. Each investor should consider whether an investment strategy of the purchase or sale of any product or security is appropriate for them in the light of their investment needs, objectives and financial circumstances.

This document constitutes a 'financial promotion' for the purposes of section 21 Financial Services and Markets Act 2000 (United Kingdom) ('FSMA') and accordingly has been approved by Capital Markets Strategy Ltd which is authorised and regulated by the Financial Conduct Authority (FCA).

No part of this document may be reproduced, stored in a retrieval system or transmitted in any form or by any means, mechanical, photocopying, recording or otherwise, without prior permission from Hardman & Co. By accepting this document, the recipient agrees to be bound by the limitations set out in this notice. This notice shall be governed and construed in accordance with English law. Hardman Research Ltd, trading as Hardman & Co, is an appointed representative of Capital Markets Strategy Ltd and is authorised and regulated by the FCA under registration number 600843. Hardman Research Ltd is registered at Companies House with number 8256259.

(Disclaimer Version 8 – Effective from August 2018)

Status of Hardman & Co's research under MiFID

Hardman & Co's research is paid for by the companies, legal entities and issuers about which we write and, as such, falls within the scope of 'acceptable minor non-monetary benefits', as defined in the Markets in Financial Instruments Directive.

<u>The FCA Handbook (COBS 2.3A.19)</u> states: 'An acceptable non-monetary benefit is one which:[...] (5) consists of: [...] (b) written material from a third party that is commissioned and paid for by a corporate issuer or potential issuer to promote a new issuance by the company, or where the third party firm is contractually engaged and paid by the issuer to produce such material on an ongoing basis, provided that the relationship is clearly disclosed in the material and that the material is made available at the same time to any firms wishing to receive it, or to the general public.'

The fact that Hardman & Co is commissioned to write the research is disclosed in the disclaimer, and the research is widely available.



research@hardmanandco.com

9 Bonhill Street London EC2A 4DJ

www.hardmanandco.com