CSL Limited

2020 Full Year Results

19th August, 2020
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A strong year for CSL with revenue up 9%¹ and profit after tax up 17%¹ reflecting:

- Strong growth in immunoglobulin portfolio
- Successful evolution of Haemophilia portfolio, driven by IDELVION®
- Transitioned to own distribution model in China
- Seqirus delivers on product differentiation strategy with strong profit growth

FY20 Revenue Performance¹
A strong year for CSL

**CSL Behring**
- PRIVIGEN® +20%
- HIZENTRA® +34%
- ALBUMIN® (36%) (GSP impact)
- IDELVION® +25%
- AFSTYLA® +21%
- HAEGARDA® +12%
- KCENTRA® +12%
- ZEMAIRA® +20%

**Seqirus**
- Seasonal influenza sales +21%
- FLUAD®:
  - Preferred recommendations in UK and Australia
  - QIV launched in Australia and approved in USA & EU
- FLUCELVAX® launched EU

¹. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.
## CSL Behring Sales FY20

<table>
<thead>
<tr>
<th>Therapy</th>
<th>Sales $m</th>
<th>Change¹ %</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Immunoglobulins</strong></td>
<td>4,014</td>
<td>22%</td>
</tr>
<tr>
<td>- IVIG</td>
<td>2,699</td>
<td>16%</td>
</tr>
<tr>
<td>- SCIG</td>
<td>1,315</td>
<td>34%</td>
</tr>
<tr>
<td><strong>Albumin</strong></td>
<td>640</td>
<td>(36%)</td>
</tr>
<tr>
<td><strong>Haemophilia</strong></td>
<td>1,122</td>
<td>8%</td>
</tr>
<tr>
<td>- Recombinants</td>
<td>659</td>
<td>18%</td>
</tr>
<tr>
<td>- Plasma</td>
<td>463</td>
<td>(3%)</td>
</tr>
<tr>
<td><strong>Specialty</strong></td>
<td>1,697</td>
<td>10%</td>
</tr>
<tr>
<td>- Peri-Operative Bleeding</td>
<td>788</td>
<td>10%</td>
</tr>
<tr>
<td>- Other Specialty</td>
<td>909</td>
<td>9%</td>
</tr>
<tr>
<td><strong>Other²</strong></td>
<td>188</td>
<td>(1%)</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>7,661</td>
<td>8%</td>
</tr>
</tbody>
</table>

1. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.

2. Includes Hyperimmunes

![Region Chart](chart.png)

US$7.7B

**Region¹**

- North America 52%
- EU 29%
- Asia Pac 11%
- ROW 8%

**19%**

**29%**

**13%**

**8%**

**16%**
**Immunoglobulins**

Sales up 22%¹

20% growth¹

- CIDP indication in the US and EU
- Expansion of SID usage
- Continued growth in PID

34% growth¹

- Clear market leader
- New patient starts in PID
- Orphan exclusivity for CIDP in the US

**Market**

- Global Ig demand remains strong

**Market Demand Drivers**

- Increased disease awareness & improved diagnosis
- Increased usage for chronic therapies
- CIDP indication
- Expanding usage for SID

¹. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.
Albumin
Sales down 36%¹

• Volume up 16% globally, excluding China:
  - Europe up 24%
  - North America up 6%
  - Emerging markets, excluding China, up 28%
  - Pricing pressure in some markets

• China:
  - One-off financial impact from GSP in line with previous guidance
  - Market volume demand outlook mid to high single digits
  - Competitive environment

Transitioned to Good Supply Practices (GSP) license in China

• Successful transition of business model
• Helps build brand and expand coverage to lower tier cities and hospitals
• No impact to patient supply

¹. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.
Haemophilia
Sales up 8%\(^1\)

- 25% growth\(^1\)
- Differentiated product
- Strong growth in US, Japan and Switzerland
- Approval of 21-day dosing in EU, Switzerland and Japan

Recombinant Coags +18%\(^1\)

Plasma Derived Coagulation Factors

- Modest growth in HUMATE\(^\circledR\)/HAEMATE\(^\circledR\) (vWF)
- pdVIII competitive pressures
- MONONINE\(^\circledR\) to IDELVION\(^\circledR\) switches

PD Coags (3%)\(^1\)

1. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.
Specialty Products
Sales increased by 10%¹

• 12%¹ growth
• Deeper penetration into broad hospital segment
• Maintained global leadership position
• 5% growth

1. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.

- 12%¹ growth
- Capacity expansion
- New launches in EU and Canada
- 10%¹ growth
- 20%¹ growth
- Supply normalised
- 9% decline
- Return of competitor

US$1,697m
Plasma Collections

Continue to grow plasma collection network

40 new centres opened in the United States

277 centres:
✓ 261 United States
✓ 8 Germany
✓ 3 Hungary
✓ 5 China

Plan to open 20 - 30 new centres in FY21

ALL CENTERS REMAIN OPEN
Plasma Collections
COVID-19 Impact

**Challenge**
- Plasma collections adversely impacted
- FY20 plasma collection volume down ~5% v FY19
- Additional collection costs incurred

**Mitigation**
- Collection centres designated ‘essential critical infrastructure’
- FDA approved inventory hold reduction from 60 to 45 days
- Utilisation of available finished goods inventory
- Potential to accelerate plasma collections
  - Enhanced marketing initiatives to increase collections
  - Investing in new centres

**Actions**
- Pre-assessment of potential donors
- Re-direction of donors to sister centers if needed
- Plasma designed social distancing
- Enhanced cleaning & disinfectant procedures
- ‘Safe passage’ letters provided to staff, donors and key vendors
## Seqirus Revenue FY20

Revenue up 11%\(^1\)

<table>
<thead>
<tr>
<th>Therapy</th>
<th>Sales $m</th>
<th>Change(^1) %</th>
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</thead>
<tbody>
<tr>
<td>QIV</td>
<td>542</td>
<td>27%</td>
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<tr>
<td>TIV</td>
<td>31</td>
<td>(53%)</td>
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<tr>
<td>Adjuvanted</td>
<td>379</td>
<td>30%</td>
</tr>
<tr>
<td>Other / In-licence</td>
<td>184</td>
<td>(11%)</td>
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<tr>
<td><strong>Total Product Sales</strong></td>
<td><strong>1,136</strong></td>
<td><strong>14%</strong></td>
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<tr>
<td>Pandemic</td>
<td>145</td>
<td>11%</td>
</tr>
<tr>
<td>Other Income</td>
<td>16</td>
<td>(64%)</td>
</tr>
<tr>
<td><strong>Total Revenue</strong></td>
<td><strong>1,297</strong></td>
<td><strong>11%</strong></td>
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</tbody>
</table>

1. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.

### Regional Sales

- **US$1.3B**
  - **57%** North America
  - **19%** Europe
  - **17%** Asia Pacific
  - **2%** ROW

- Seasonal Influenza vaccines +21%

- **18%**
  - **11%** US
  - **11%** Other Regions

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Driven by **Our Promise™**
Operating Highlights

• Strong result driven by ongoing product differentiation

• Critical operations maintained during COVID pandemic

• Real world evidence continues to demonstrate the potential for improved effectiveness of FLUCELVAX® & FLUAD®

• Progression on fill and finish expansion projects

Looking Forward

• COVID-19 driving demand:
  – Increased supply into the US of up to ~60m doses for NH20/21

• Fill & finish expansion:
  – Liverpool operational from NH 21/22
  – Holly Springs operational from NH 22/23
Seqirus

Operating Highlights

FLUCELVAX®
- All strains manufactured using cell-specific seed for NH 2019/20 season
- Launched in EU (9 years+)

FLUAD®
- Ongoing preferred recommendations in UK and Australia
- 65+ QIV launched in Australia; approved in US, EU and UK

AFLURIA®
- QIV approved in Argentina and Germany

Looking Forward

FLUCELVAX®:
- Expanded paediatric in US and EU 2021
- aQIVc to commence clinical trials NH20/21

FLUAD QIV®:
- US NH20/21 and EU NH21/22 launches
R&D Highlights

**Immunology**
- HIZENTRA® Phase III DM study initiated
- PRIVIGEN® Phase II SSc study initiated
- HAEGARDA® Phase III HAE study in Japan initiated
- PRIVIGEN® approved for PID & SSc in Japan
- Garadacimab Phase II HAE study results presented at EAACI Congress
- FDA granted HIZENTRA® orphan drug exclusivity for CIDP; PRIVIGEN® ODD and fast track designation for SSc
- Alliance with Seattle Children’s Research Institute to develop stem cell gene therapies for PID – WAS and XLA

**Hematology**
- CSL200 in SCD Phase I study initiated
- CSLB89 Hemopexin Phase I SCD study initiated
- FDA granted CSL200 fast track designation
- CSLB89 Hemopexin ODD approved in EU for SCD
- CSL agreed to acquire exclusive global license rights to adeno-associated virus gene therapy program, AMT-061 EtranaDez for hemophilia B*

*The transaction with uniQure is subject to customary regulatory clearances before closing

**Cardiovascular & Metabolic**
- CSL112 (ApoA-I) Phase III study (AEGIS-II) >9500 patients recruited
- CSL112 AEGIS-II first futility analysis conducted; trial to continue as planned

**Transplant**
- AAT for prevention of GvHD Phase III study enrolment into Cohort 2 completed
- CSL acquired Vitaeris Inc. and Clazakizumab
- Clazakizumab AMR study initiated
- FDA granted Clazakizumab ODD and fast track designation

**Influenza Vaccines**
- First cell-based quadrivalent seasonal influenza vaccine, FLUCELVAX® TETRA, approval in Europe
- US FDA approval of AUDENZTM - adjuvanted, cell-based influenza A (H5N1) pandemic vaccine
- aQVC (cell antigen + MF59) new product development commenced
COVID-19 Response

### Therapeutic Options

**Vaccines**

- University of Queensland
- Coalition for Epidemic Preparedness Innovations (CEPI)

**Hyperimmunes Polyclonal Antibodies**

- Hyperimmunes
  - Alliance with Takeda and others
  - Australian program
- Polyclonal: SAB Therapeutics

**Monoclonal Antibodies**

- Academic clinical researchers

### Collaborators

- Partnership formed to accelerate the development, manufacture and distribution of vaccine
- Vaccine to be available in 2021 if successful

### Update

- FPI achieved for first mAb offering (CSL312)
- CSL324 IND submitted June 20; FPI expected in 3Q20

<table>
<thead>
<tr>
<th>PREVENTION</th>
<th>TREATMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vaccines</strong></td>
<td><strong>Hyperimmunes Polyclonal Antibodies</strong></td>
</tr>
</tbody>
</table>
| University of Queensland | Hyperimmunes
- Alliance with Takeda and others
- Australian program | Academic clinical researchers |
| Coalition for Epidemic Preparedness Innovations (CEPI) | Polyclonal: SAB Therapeutics | |
| Partnership formed to accelerate the development, manufacture and distribution of vaccine | Clinical manufacture underway | |
| Vaccine to be available in 2021 if successful | Clinical trial start targeted for this quarter | |

Driven by Our Promise™
COVID-19 Summary

COVID-19 presents some challenges however we remained focussed on strategy execution and continue to invest for growth.

**PEOPLE**
- Recognised as an ‘essential’ business
- Employees encouraged and supported to work remotely
- Flexible and robust IT systems facilitate ongoing productivity
- Development of strict protocols to ensure the safety of our employees and donors

**INNOVATION**
- Focussed response leveraging the Company capabilities:
- Prevention – vaccine collaboration
- Treatment:
  - Hyperimmune – Global, Australia and SAB
  - Pivoting Mabs and plasma products into ARDS patients
- Paused clinical trials to recommence in FY21

**DEMAND**
- Products are used to treat serious rare diseases and often used chronically
- Demand remains strong across the portfolio
- Especially strong for IG & influenza vaccines
- Increased preference for home treatment driving HIZENTRA® demand

**SUPPLY**
- All plasma centres remain open
- CSL Behring & Seqirus manufacturing facilities operational
- Plasma collections adversely impacted by COVID-19 pandemic
- Multiple initiatives underway to ensure patient supply of therapies

**BALANCE SHEET**
- Ongoing conservative approach to liquidity and leverage
- Raised US$750 million via private placement, bolstering existing strong capital position
- Net debt to EBITDA 1.5x. Available liquidity $3.1 billion
- Credit ratings S&P A-, Moody’s A3
Financials

David Lamont
CFO
Financial Highlights
Net Profit After Tax

<table>
<thead>
<tr>
<th>FY19</th>
<th>FY20</th>
</tr>
</thead>
<tbody>
<tr>
<td>$1,919 m</td>
<td>$2,247m @CC¹</td>
</tr>
<tr>
<td>+17% @CC¹</td>
<td>FX$144m</td>
</tr>
<tr>
<td>+10% reported</td>
<td></td>
</tr>
</tbody>
</table>

1. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.

GSP China Transition
• Albumin sales reduction in line with guidance
• Profit effect in line with historical CSL Behring margin

Other Income
• One-off $30m benefit

New lease standard
• Balance sheet gross up
• P&L impact immaterial
# Financial Highlights

## CSL Group

<table>
<thead>
<tr>
<th>Financial Highlight</th>
<th>FY19 Reported</th>
<th>FY20 Reported</th>
<th>FY20 at CC</th>
<th>Change %</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Revenue</strong></td>
<td>8,539</td>
<td>9,151</td>
<td>9,295</td>
<td>9%</td>
</tr>
<tr>
<td><strong>Gross Profit</strong></td>
<td>4,777</td>
<td>5,226</td>
<td>5,338</td>
<td>12%</td>
</tr>
<tr>
<td><strong>GP margin</strong></td>
<td>56.0%</td>
<td>57.1%</td>
<td>57.4%</td>
<td></td>
</tr>
<tr>
<td><strong>EBIT</strong></td>
<td>2,504</td>
<td>2,717</td>
<td>2,877</td>
<td>15%</td>
</tr>
<tr>
<td><strong>EBIT margin</strong></td>
<td>29.3%</td>
<td>29.7%</td>
<td>31.0%</td>
<td></td>
</tr>
<tr>
<td><strong>NPAT</strong></td>
<td>1,919</td>
<td>2,103</td>
<td>2,247</td>
<td>17%</td>
</tr>
<tr>
<td><strong>Cashflow from Operations</strong></td>
<td>1,644</td>
<td>2,488</td>
<td></td>
<td>51%</td>
</tr>
<tr>
<td><strong>ROIC</strong></td>
<td>24.3%</td>
<td>21.6%</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>EPS ($)</strong></td>
<td>4.24</td>
<td>4.63</td>
<td>4.95</td>
<td>17%</td>
</tr>
<tr>
<td><strong>DPS ($)</strong></td>
<td>1.85</td>
<td>2.02</td>
<td></td>
<td>9%</td>
</tr>
</tbody>
</table>

1. Constant Currency (CC) removes the impact of exchange rates movements to facilitate comparability. See end note for further detail.
# Financial Highlights

## Segments

### CSL Behring

<table>
<thead>
<tr>
<th>US$ Millions</th>
<th>FY19 Reported</th>
<th>FY20 Reported</th>
<th>Change % at CC¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sales</td>
<td>7,187</td>
<td>7,661</td>
<td>8%</td>
</tr>
<tr>
<td>Other Revenue</td>
<td>156</td>
<td>193</td>
<td>24%</td>
</tr>
<tr>
<td>Total Revenue</td>
<td>7,343</td>
<td>7,854</td>
<td>9%</td>
</tr>
<tr>
<td>Gross Profit</td>
<td>4,195</td>
<td>4,540</td>
<td>11%</td>
</tr>
<tr>
<td>GP margin</td>
<td>57.1%</td>
<td>57.8%</td>
<td></td>
</tr>
<tr>
<td>EBIT</td>
<td>2,351</td>
<td>2,451</td>
<td>11%</td>
</tr>
<tr>
<td>EBIT margin</td>
<td>32.0%</td>
<td>31.2%</td>
<td></td>
</tr>
</tbody>
</table>

### Seqirus

<table>
<thead>
<tr>
<th>US$ Millions</th>
<th>FY19 Reported</th>
<th>FY20 Reported</th>
<th>Change % at CC¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sales</td>
<td>1,018</td>
<td>1,136</td>
<td>14%</td>
</tr>
<tr>
<td>Other Revenue</td>
<td>178</td>
<td>161</td>
<td>(7%)</td>
</tr>
<tr>
<td>Total Revenue</td>
<td>1,196</td>
<td>1,297</td>
<td>11%</td>
</tr>
<tr>
<td>Gross Profit</td>
<td>582</td>
<td>686</td>
<td>17%</td>
</tr>
<tr>
<td>GP margin</td>
<td>48.7%</td>
<td>52.9%</td>
<td></td>
</tr>
<tr>
<td>EBIT</td>
<td>153</td>
<td>265</td>
<td>74%</td>
</tr>
<tr>
<td>EBIT margin</td>
<td>12.8%</td>
<td>20.4%</td>
<td></td>
</tr>
</tbody>
</table>

¹. Growth percentages shown at constant currency to remove the impact of exchange rate movements, facilitating comparability of operational performance. See end note for further detail.
Financial Highlights

Reported Expenses

<table>
<thead>
<tr>
<th></th>
<th>FY20 $m</th>
<th>Change @ CC(^1) $m</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Research &amp; Development</td>
<td>922</td>
<td>101</td>
<td>12</td>
</tr>
<tr>
<td>Sales &amp; Marketing</td>
<td>896</td>
<td>39</td>
<td>5</td>
</tr>
<tr>
<td>General &amp; Admin</td>
<td>692</td>
<td>48</td>
<td>8</td>
</tr>
</tbody>
</table>

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Finance (Net)</td>
<td>144</td>
<td>(28)</td>
<td>(17)</td>
</tr>
<tr>
<td>Tax</td>
<td>470</td>
<td>72</td>
<td>17</td>
</tr>
<tr>
<td><strong>ETR %</strong></td>
<td></td>
<td></td>
<td>18.3%</td>
</tr>
</tbody>
</table>

R&D
- CSL112 phase III trial
- CSL200 SCD gene therapy trial initiated
- COVID-19 response

Finance (Net)
- AASB16 adoption - $26m
- YoY variation in Swiss debt costs $41m

Tax
- Increase in-line with profit increase
- FY21 ETR rate ~20-22%

1. Constant Currency (CC) removes the impact of exchange rates movements to facilitate comparability. See end note for further detail
Inventory Management

Flexibility in the supply chain

Key Insights

• Continue to produce for demand
• Inventory as a percentage of revenue steady
• Seqirus inventory mix impacted by strain notification

1 Reported numbers
Capital Expenditure\(^1\)
Investment to support demand

Key Projects
- Significant new manufacturing capacity
- Additional plasma collection centers
- ERP systems completed
- Lengnau facility
  - Thermo Fisher lease
  - Accelerates optimisation of facility capability
- FY21 ~ $1.6 billion

\(^1\) Reported numbers
## Key Capital Projects

### Completion Timeline

<table>
<thead>
<tr>
<th>Tech</th>
<th>FY20</th>
<th>FY21</th>
<th>FY22</th>
<th>FY23</th>
<th>FY24</th>
<th>FY25</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERP Systems</td>
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<td>Enterprise Process Management</td>
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<tr>
<td>Kankakee</td>
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<td>Base Frac</td>
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<tr>
<td>Marburg</td>
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<td>Broadmeadows</td>
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<tr>
<td>Other Base Frac Modules</td>
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<td>Bulk &amp; Finishing</td>
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<td>Bern IG</td>
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<td>IG Modules</td>
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<td>Berinert / Cl Precipitate Capacity Increase</td>
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<td>Other</td>
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<td>Plasma Centers (Continuous)</td>
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<td>Other</td>
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<td>Seqirus</td>
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<td>Biotech Facility</td>
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<td>Fill &amp; Finish – Liverpool</td>
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<td>Fill &amp; Finish – Holly Springs</td>
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Driven by Our Promise™
Outlook for FY21

Demand
• Continued strong demand for plasma and recombinant products
• Seqirus’ product differentiation and COVID-19 expected to drive strong demand for influenza vaccines
• Albumin sales to normalize following GSP transition

Plasma Collections
• COVID-19 restrictions expected to restrain plasma collections
• Additional plasma collection costs
• Multiple initiatives underway to mitigate impact

R&D
• COVID-19 response and new growth initiatives to drive uplift in investment towards the top end of prior guidance range

FY21 Outlook

Revenue Growth
~6 - 10% @CC

NPAT
~$2,100 - $2,265m @CC

1 For forward looking statements, refer to Legal Notice on page 2.
2 Constant Currency (CC) removes the impact of exchange rates movements to facilitate comparability. See end note for further detail.
3 Previously provided R&D investment guidance of ~10-11% of revenue.
CSL Contacts

Mark Dehring  
VP Investor Relations  
📞 +61 3 9389 3407  
✉️ mark.dehring@csl.com.au

Bernard Ronchi  
Senior Manager, Investor Relations  
📞 +61 3 9389 3470  
✉️ bernard.ronchi@csl.com.au

Stephen McKeon  
Associate Director, Investor Relations  
📞 +61 3 9389 6798  
✉️ stephen.mckeon@csl.com.au
CSL Strategy and Values

Strategic Overview

Core Values

- **Patient Focus**: We deliver on our promise to patients
- **Innovation**: We turn innovative thinking into solutions
- **Integrity**: We walk the talk
- **Collaboration**: We are stronger together
- **Superior Performance**: We take pride in our results
Notes

[1] Constant currency removes the impact of exchange rate movements to facilitate comparability of operational performance for the Group. This is done in three parts: a) by converting the current year net profit of entities in the group that have reporting currencies other than US Dollars, at the rates that were applicable to the prior comparable period (translation currency effect); b) by restating material transactions booked by the group that are impacted by exchange rate movements at the rate that would have applied to the transaction if it had occurred in the prior comparable period (transaction currency effect); and c) by adjusting for current year foreign currency gains and losses. The sum of translation currency effect, transaction currency effect and foreign currency gains and losses is the amount by which reported net profit is adjusted to calculate the operational result.

Summary NPAT
Reported net profit after tax $2,102.5m
Translation currency effect (a) $ (1.0m)
Transaction currency effect (b) $ 60.1m
Foreign Currency (gains) & losses (c) $ 85.4m
Constant currency net profit after tax * $2,247.0m

a) Translation Currency Effect $(1.0m)
Average Exchange rates used for calculation in major currencies (Twelve months to June 20/June 19) were as follows: USD/EUR (0.90/0.87); USD/CHF (0.98/0.99).

b) Transaction Currency Effect $60.1m
Transaction currency effect is calculated by reference to the applicable prior year exchange rates. The calculation takes into account the timing of sales both internally within the CSL Group (ie from a manufacturer to a distributor) and externally (ie to the final customer) and the relevant exchange rates applicable to each transaction.

c) Foreign Currency Loss $85.4m
Foreign currency gains recorded during the period.

Summary Sales
Reported sales $8,796.6m
Currency effect $ 141.2m
Constant currency sales* $ 8,937.8m

* Constant currency net profit after tax and constant currency sales have not been audited or reviewed in accordance with Australian Auditing Standards.