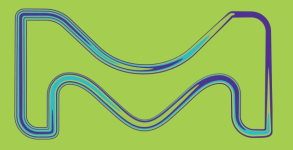
Merck KGaA, Darmstadt, Germany 101 21 Roadshow

Peter Guenter, CEO Healthcare Marcus Kuhnert, CFO

May 2021





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Risks and uncertainties include, but are not limited to: the risks of more restrictive regulatory requirements regarding drug pricing, reimbursement and approval; the risk of stricter regulations for the manufacture, testing and marketing of products; the risk of destabilization of political systems and the establishment of trade barriers; the risk of a changing marketing environment for multiple sclerosis products in the European Union; the risk of greater competitive pressure due to biosimilars; the risks of research and development; the risks of discontinuing development projects and regulatory approval of developed medicines; the risk of a temporary ban on products/production facilities or of non-registration of products due to non-compliance with quality standards; the risk of an import ban on products to the United States due to an FDA warning letter; the risks of dependency on suppliers; risks due to product-related crime and espionage; risks in relation to the use of financial instruments; liquidity risks; counterparty risks; market risks; risks of impairment on balance sheet items; risks from pension obligations; risks from product-related and patent law disputes; risks from antitrust law proceedings; risks from drug pricing by the divested Generics Group; risks in human resources; risks from e-crime and cyber attacks; risks due to failure of business-critical information technology applications or to failure of data center capacity; environmental and safety risks; unanticipated contract or regulatory issues; a potential downgrade in the rating of the indebtedness of Merck KGaA, Darmstadt, Germany; downward pressure on the common stock price of Merck KGaA, Darmstadt, Germany and its impact on goodwill impairment evaluations as well as the impact of future regulatory or legislative actions.

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Agenda

- **Business overview**
- **O2** Transforming the company
- Healthcare Executing on the earnings phase
- Life Science Focusing on profitable growth
- **Electronics Leveraging portfolio shift**
- **Sustainability**
- **O**7 **Guidance & executive summary**



Business overview

Ol

Three high-tech businesses competing in attractive markets



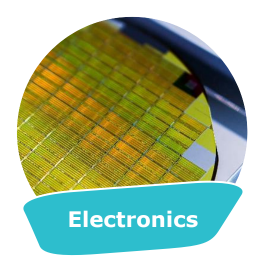
Leading in specialty pharma markets

- Biologics and small-molecule prescription medicines against cancer, multiple sclerosis, infertility
- Research focus: Oncology,
 Immunology & Immuno-Oncology
- Successful portfolio management: e.g. divestment of Consumer Health and Allergopharma



Leading life science company

- Tools and services for biotech research & production
- Tools and laboratory supply for academic research and industrial testing



Leading company in high-tech solutions

- High-tech solutions and materials for electronics
- BroaMattd portfolio of decorative and functional solutions



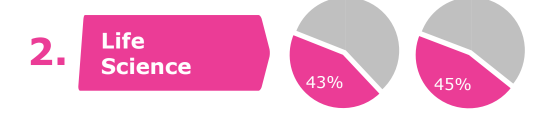
Group today – three strong pillars as basis for profitable growth





Global specialty innovator poised for above-industry growth

- Resilient core business backed by excellent life cycle management
- Strong growth from new products, late-stage pipeline assets with blockbuster potential
- Rigorous cost discipline and value-maximizing pipeline prioritization



Diversified industry leader poised for above-market growth

- Portfolio advantage and outperformance drive above-market growth
- Strengthen core: products (PS), chemistry (RS), lab water (AS)
- Establish new pillars: PS services, gene editing and novel modalities



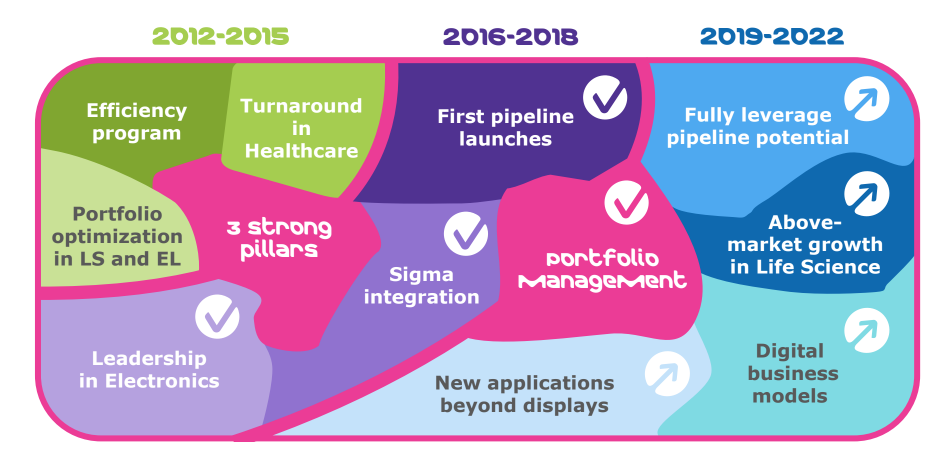
Leading electronics player poised for accelerating growth

- Growing semiconductor share as key driver for acceleration
- More resilient growth through rising diversification
- Strict cost discipline in maturing parts of the portfolio

¹EBITDA pre share excluding Corporate & Others; 2020 EBITDA pre restated for €365 m patent litigation provision release Acronyms: PS = Process Solutions, RS = Research Solutions, AS = Applied Solutions



The 2016 vision – a strategic agenda until 2022



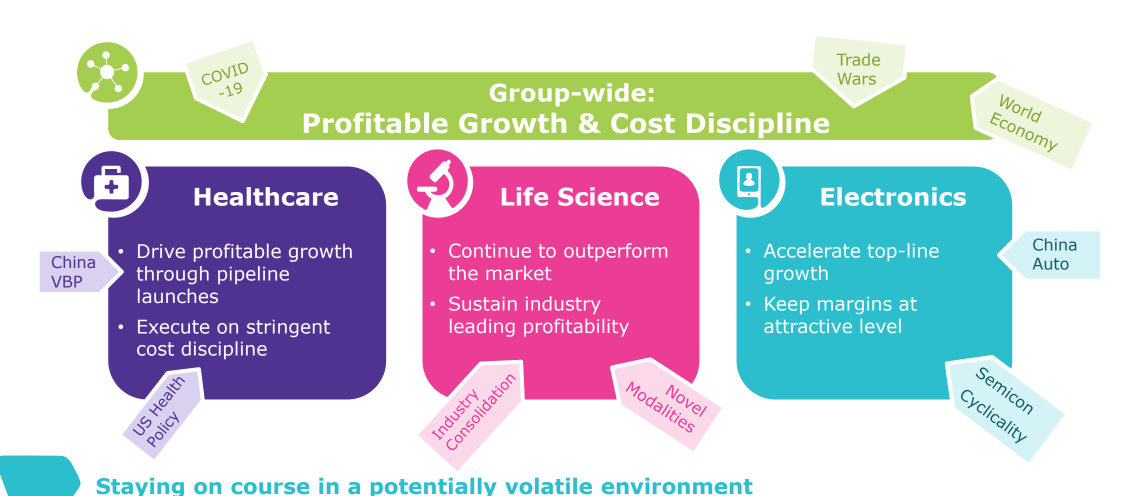


Executing on the growth and expansion phase of the 2016-22 strategic agenda





2021 and beyond - poised for growth in a challenging environment



Acronym: VBP = volume based procurement



the company

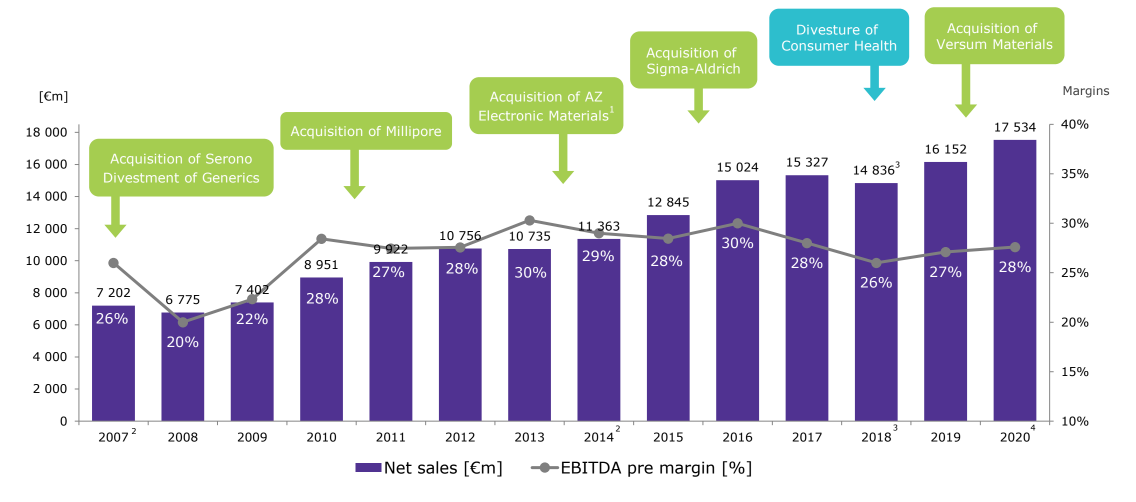
02

We have added scale and strengthened the attractiveness of our portfolio





Continue to transform to a science and technology focused company



¹Included since 2 May 2014; ²2007 and 2014 EBITDA pre margin adjusted for comparability; ³2018 net sales reflect Consumer Health divesture (reduction of ~€1 bn net sales p.a.) ⁴2020 margin restated for €365 m patent litigation provision release



All three business sectors delivering on their strategic priorities

Healthcare

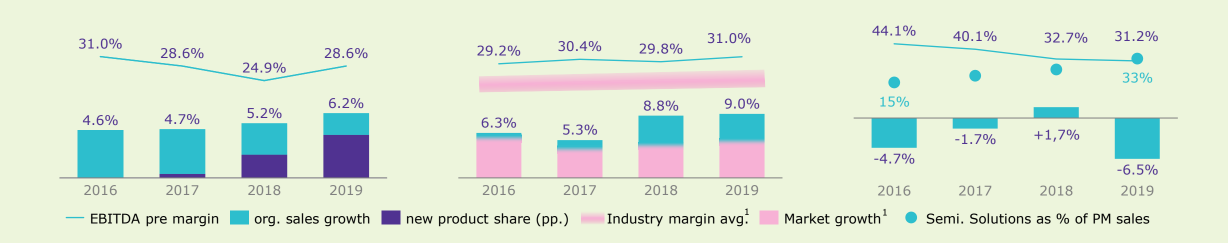
- Accelerating organic growth with rising contribution from launches (Mavenclad[®], Bavencio[®])
- Margin trough behind, pipeline progressing well

Life Science

- Above-market organic growth paired with industry-leading margin
- Significant growth investments (organic, inorganic & partnerships)

Electronics

- Significant portfolio change towards higher growth business (Semi, OLED)
- Margin stabilizing at ~30% amid sound execution of Bright Future

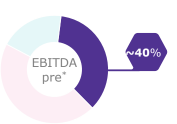


¹ Company estimate based on industry data and reporting by peers

Clear set of priority goals



Healthcare



- Deliver on ambition to keep core business at least stable until 2022
- Transition from investment to earnings phase by 2019
- Foster successful Bavencio[®]
 and Mavenclad[®] ramp up
- Stringent pipeline execution



Life science



- Strengthen position as differentiated player in a highly attractive market
- Maintain consistent abovemarket growth trajectory and superior profitability





- Deliver ambition of 3-4% CAGR
- Implement 5-year transformation program and focus on seamless integration of Versum and Intermolecular; to deliver financial ambition of around 30% margin
- Maintain strong cash generation and cash conversion



*based on FY 2020

Three-pillar structure – positioned to win in high-growth markets

Global economy¹



Global **GDP**

~3% to 4%





Global pharma industry ~4% to 5%



Global life science industry



Global electronics industry ~4%



~4% to 5%

Focus market areas¹



Oncology: ~10%

Immunology: ~5% to 9%



Biologics: ~10% to 12%

Services: ~7% to 8%



Semi materials: ~4% to 6%

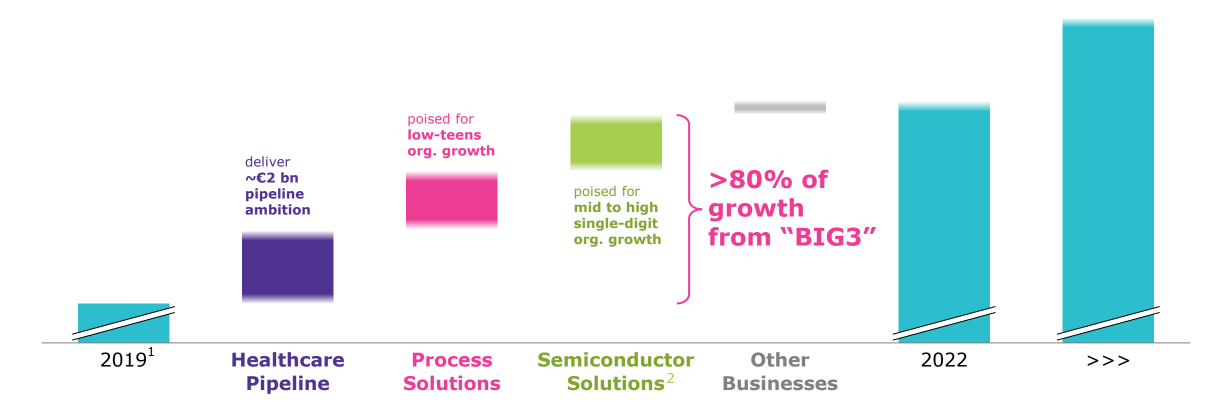




Purposefully positioned in attractive markets with secular growth above global GDP ...further focusing investments on attractive sub-segments

¹ Company estimates of mid-term growth outlook based on industry forecasts and reports from public research institutes (e.g. IMF, IQVIA, EvaluatePharma, Prismark, etc.)

Three main drivers of growth to 2022 and beyond



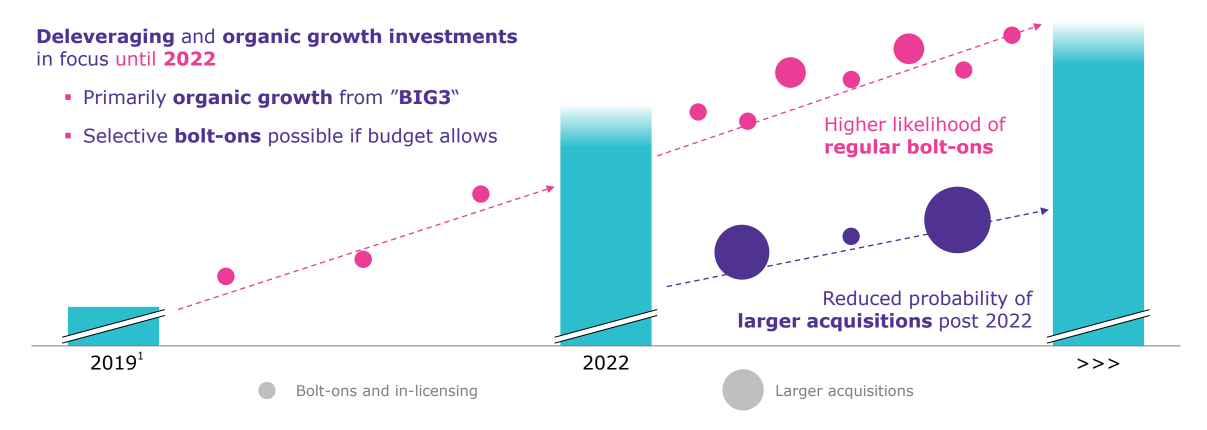


Beyond 2022: further significant growth potential from "BIG3" and increasing contributions from other businesses

¹ 2019 Group sales of €16.2 bn; ² Including Versum portfolio effect



Portfolio strategy - from transformation to evolution





Strong portfolio: significant organic growth potential to 2022 and beyond ...and higher likelihood of regular bolt-ons post 2022

¹ 2019 Group sales of €16.2 bn

Healthcare

Executing on the earnings phase

03

Healthcare

Creating optionality through focused pipeline approach

Pipeline and launch progress supported by strong base

Pipeline quality

- Early stage pipeline fueled by in-house innovation
- Potentially **transformative late-stage pipeline** assets (Evobrutinib, Bintrafusp alfa)
- **Optimized risk profile** through strategic partnerships (Pfizer and GSK), in-licensing (pivotal-stage Xevinapant) & focused externalization

Launch activity

- Mavenclad[®] approved in >80 countries including the U.S.
- Bavencio[®] launched in MCC¹, RCC², and UC 1L/2L³
- **Tepotinib** first-in-class approval of an oral MET inhibitor •

Strong core business due to excellent life cycle management

Mid-term outlook Healthcare

~€2 bn
pipeline sales
& stable base business
until 2022
+
significant growth
potential
beyond



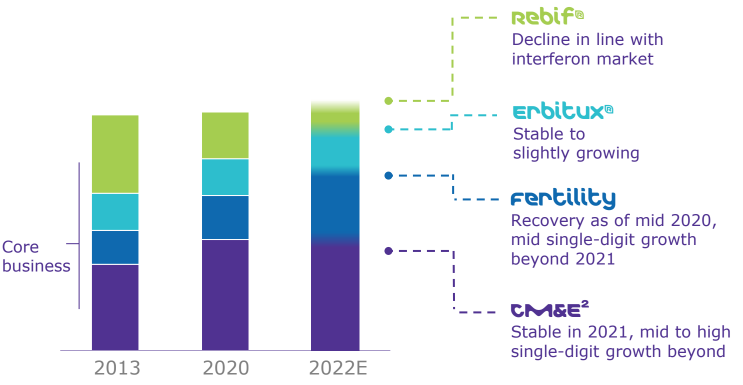
Confirming financial ambition of a stable base business and
~€2 bn pipeline sales until 2022 with further significant growth potential beyond

¹ MCC = Merkel Cell Carcinoma, launched in all major jurisdictions; ² RCC = Renal Cell Carcinoma, launched in all major jurisdictions; ³ UC = Urothelial Carcinoma, 1L = first line, 2L = second line, launched in all major jurisdictions; ⁴ approved in Japan and U.S.

Healthcare: Base Business

Confirming ambition to keep base business at least stable to 2022

Healthcare base business net sales until 2022



- Maintain solid track record of patient retention
- Integrate into joint franchise with Mavenclad®
- Drive EM¹ growth and mitigate competitive / price pressure in EU by clear branding
- Continued China NRDL inclusion secured through successful renegotiation in late 2020
- Drug demand driven by emerging markets growth and demographics
- Leverage competitive strengths (e.g. broad and innovative portfolio, security of supply)
- Drive recovery after COVID-19
- Increasing prevalence of diabetes and cardiovascular diseases
- Mitigate VBP pressure in China through EM growth, effective life cycle management, and portfolio expansion



Core business with **36 consecutive quarters of growth** (Q2 2011 – Q1 2020) Growth to **pick up after COVID-19 impact**, further **growth potential after 2022**



¹ EM: emerging markets; ² Cardiovascular, Metabolism and Endocrinology (new Franchise name as of Q1 2021)

Healthcare: Sales from Pipeline

Mavenclad and Bavencio launches on track for ~€2 bn pipeline

ambition in 2022



- Global peak sales: €1.0-1.4 bn
- Approved in >80 countries, including USA, EU, Canada and Australia
- Dynamic market volume still lags pre pandemic levels by ~20% but increased vaccination likely to drive market recovery and Q2 growth



- Approved for aRCC (USA, EU, Japan), mMCC (50 countries incl. USA and EU), and UC 2L (USA, Canada, Israel)
- Approved for UC 1L in US on June 30, 2020, in EU on January 25, 2021 and in Japan on February 24, 2021
- Phase III read-out remaining: NSCLC 1L (est. primary completion date: October 2021)



Tepotinib

- Approved in Japan on March 25, 2020 (Sakigake and ODD granted in 2018 & 2019)
- Approved in US on February 3, 2021 (granted priority review under RTOR)

Bintrafusp alfa

 Multiple potentially registrational studies across various tumor types ongoing







Independent real-world data (RWD) differentiates Mavenclad®

- A high-efficacy DMT that demonstrates full antibody response to COVID-19 vaccination
- Differentiated vs. other high-efficacy therapies in light of COVID-19 vaccinations for MS patients

Patient population		Total N=125	Protective humoral immunity ^a
DMT treated patients	Mavenclad®	23	100% (p = 0.99) ^b
	Ocrelizumab	44	22.7% (<i>p</i> < 0.0001) ^b
	Fingolimod	26	3.8% (<i>p</i> < 0.0001) ^b
Untreated MS patients		32	100%
Healthy subjects		47	97.9%



In the first-ever real-world data study of its type all patients on Mavenclad® who received a mRNA COVID-19 vaccine were able to mount a full antibody response, similar to healthy subjects and untreated people with MS, irrespective of lymphocyte counts¹

^aProtective humoral immunity defined as a index value higher than 1.1 using EUROIMMUN semiquantitiative ELISA for IgG specific for the recombinant S1 subunit of SARS-CoV-2 spike protein bFisher's exact test to detect differences in categorical variables between DMT-treated patients with MS and untreated patients with MS



DMT = disease-modifying therapy

^{1.}Achiron et al. Ther Adv Neurol Disord https://doi.org/10.1177/17562864211012835

Bavencio® UC 1L launch update: Significant opportunity to drive further growth by increasing the adoption of 1L maintenance therapy



U.S. - ~10 months into launch, on track to changing the SOC:





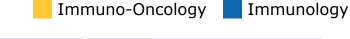


Europe & Japan - Recently approved, promising early signals:

- Market access on track
- Strong initial uptake in key launch markets
- Recently recommended by EAU (European Association of Urology) guidelines (March 25) as the preferred treatment in 1L UC



Healthcare catalysts – Significant developmental progress across Oncology and Immuno-Oncology portfolio expected in 2021



 Q2 2021
 Q3 2021
 Q4 2021
 Q1 2022

 Tepotinib (c-Met-inhibitor)

EMA: Filing accepted in Q4 2020, review ongoing

Bavencio® (Avelumab/Anti-PD-L1)

1L NSCLC (JAVELIN 100): Expected data read-out¹ Bintrafusp alfa (TGFbeta trap/anti-PD-L1)

2L Cervical (INTR@PID 017): Expected data read-out

Xevinapant (IAP inhibitor)

Initiation of Ph III study in Cisplatin-Ineligible LA SCCHN

M5049 (TLR 7/8 antagonist)

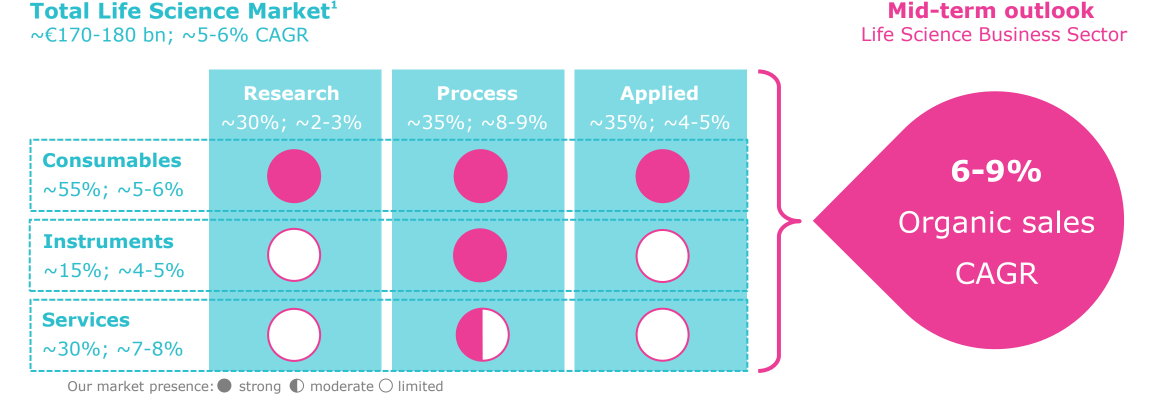
Covid-19 pneumonia: Results dependent on recruitment and COVID-19 infection rates



Focusing on profitable growth



Building growth momentum with focus on attractive market segments

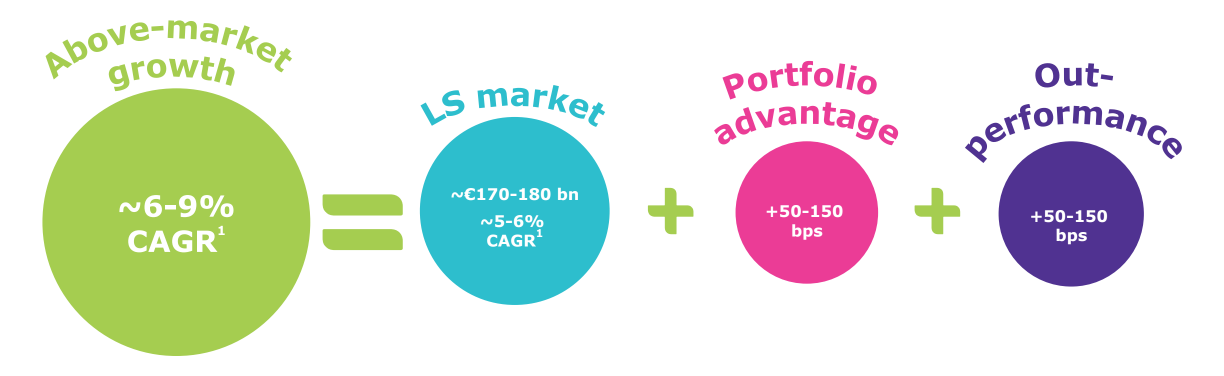




Upgrading mid-term financial ambition to 6-9% organic sales CAGR

¹ Company estimate of the market segments, based on industry forecast over 5-year horizon; all growth rates in 3x3 Matrix indicate external market growth

Improved mid-term outlook driven by market and portfolio focus

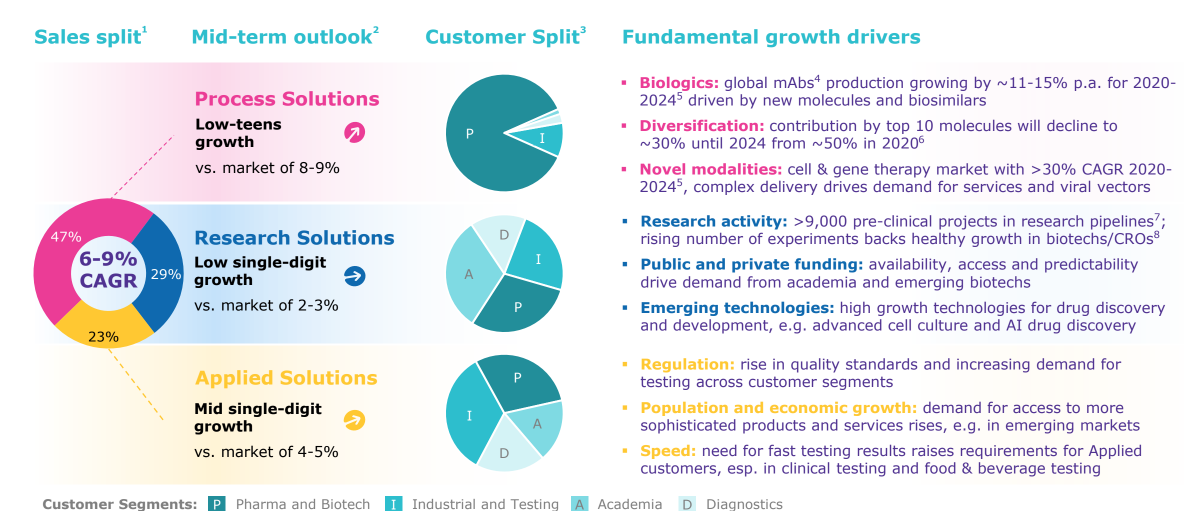






¹ Company estimate based on industry forecast over 5-year horizon

All business units contributing to above-market growth



¹ Based on H1 2020, CAGR is organic mid-term ambition; ² growth rates are organic CAGRs; ³ indicative only; ⁴ mAbs = monoclonal antibodies; ⁵ Source: company estimate based on industry forecasts; ⁶ Source: EvaluatePharma; ⁷ Source: statista; ⁸ CRO = Contract Research Organization

Critical offering in the fight against COVID-19



products feed into...

www.sigmaaldrich.com/covid-19

VIRUS **DETECTION**

- Leading critical component provider for Molecular and Serological diagnostic kits
- PCR reagents, kits and tools for all stages of assay development

VIRUS CHARACTERIZATION

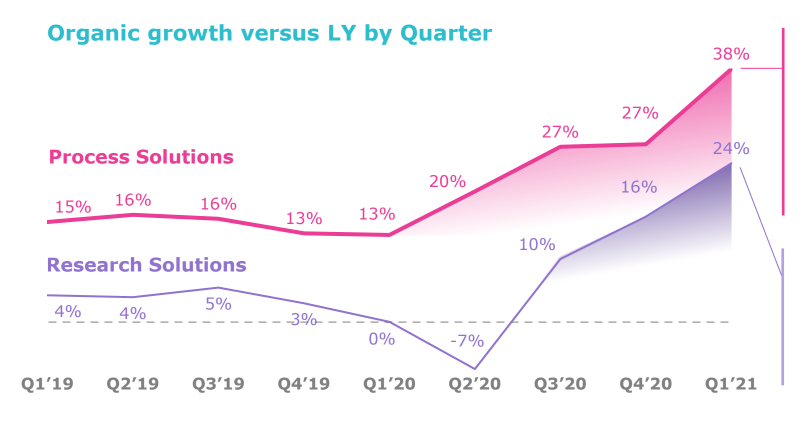
- Offering is among largest biologic reagents and hardware portfolios
- Effective vaccines and therapies start with reliable virus characterization
- Highest quality reagents needed for understanding of viral attachment, genomics, or proteomics

VACCINE & THERAPY **PRODUCTION**

- Supporting global COVID-19 vaccine and therapy response effort:
 - Upstream and downstream research and scaling
 - End-to-End solutions
 - Biosafety Testing Services



Life Science: Upside potential for Process Solutions materializing amid increasing capacity; Research Solutions gaining momentum as well



- Successful capacity ramp-up accelerated further
- Roughly half of additional growth since Q2 2020 has been COVID-19 related
- Strong underlying demand
- Order intake > +60%
- Strong core business
- Parts of growth attributable to ongoing recovery post lock-downs
- Less than half of growth since Q2
 2020 has been COVID-19 related



Key factors for 2021 guidance remain:

- Further progress of capacity expansions & optimizations
- Sustainable demand growth; both Covid-19 and underlying

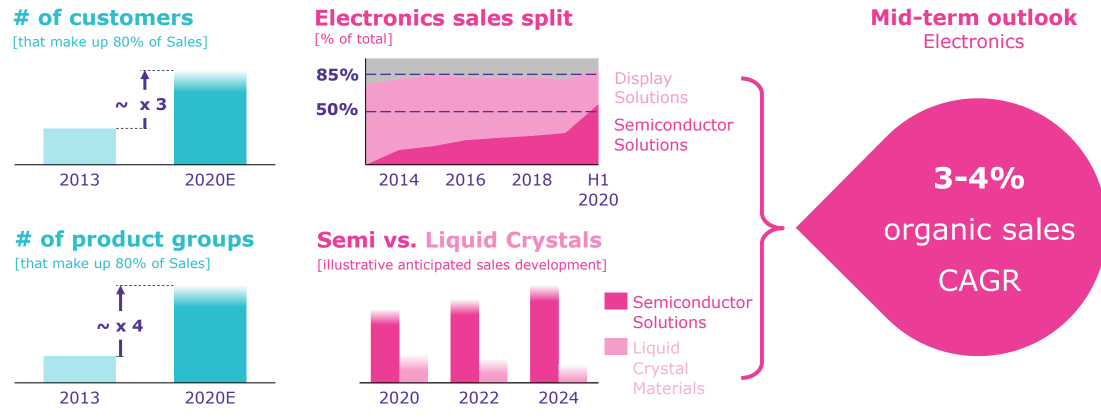




Leveraging the portfolio shift



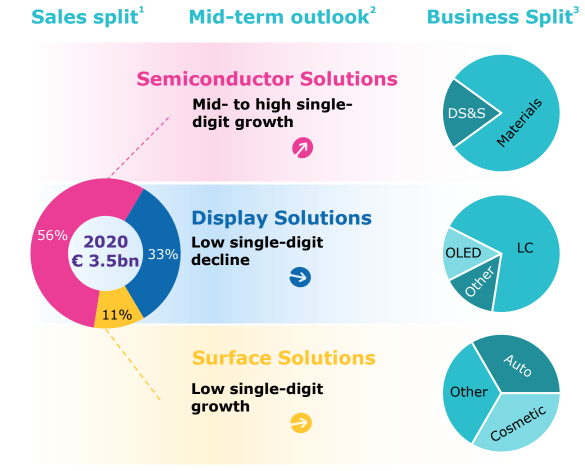
Portfolio shift leads to greater resilience and accelerated growth





Updating mid-term financial ambition to 3-4% organic sales CAGR

Portfolio refocus drives mid-term guidance upgrade to 3 to 4% CAGR



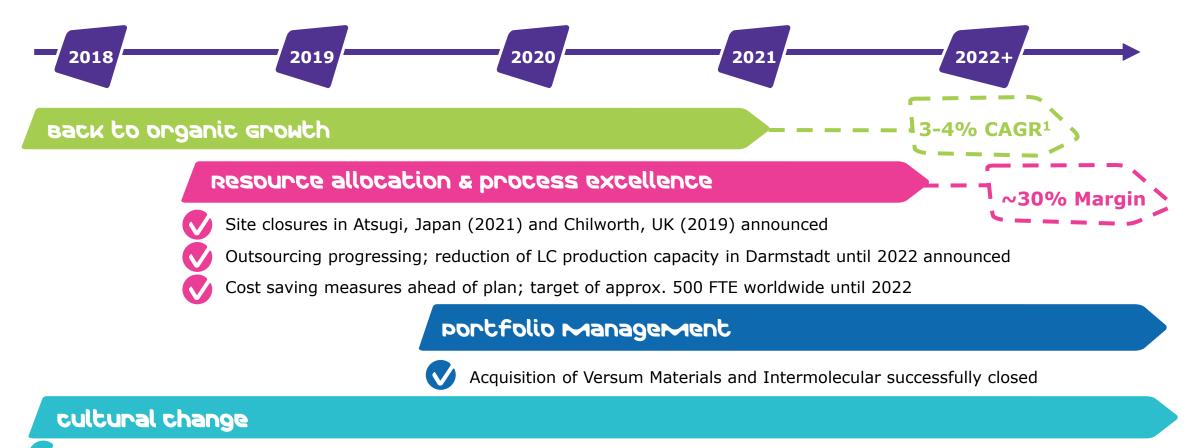
Fundamental growth drivers

- Continued market growth due to technological advances (Artificial intelligence, 5G, Big Data and cloud, Internet of Things) serving customers in Logic, Memory, Packaging and others
- 4 to 6% market growth⁴
- 200 to 300bps above-market growth from share gains & better portfolio (incl. 100 to 150bps additional growth from integration top-line synergies)
- Driven by trend to bigger TV size, higher resolutions, more mobile devices
- 3 to 4% growth of total LCD m² area⁵, while price pressure continues
- 18 to 22% growth of total OLED m² area⁵ with slight to moderate market share gains
- OLED material market to exceed LC material market by 2021⁶
- Well balanced exposure to automotive and cosmetics end market
- Drivers: rising living standards, higher disposable income in growing markets & higher demand for high value products at reasonable prices
- Light vehicle production and relevant cosmetics end markets returning to growth in 2021 and reaching 2019 levels by 2022 and beyond⁷



¹ Based on FY 2020, CAGR is organic mid-term ambition; ²growth rates are organic CAGRs; ³indicative only ⁴Source: Jan 2020 IC Insights 2018-2024 CAGR for wafer starts in million units; ⁵Source: Omdia Display Market Outlook, Q1 2020; ⁶Internal Business Intelligence; ⁷Sources: LMC Automotive Light Vehicles Forecast, Aug 2020 & Euromonitor BPC (Beauty & Personal Care) Aug 2020

5-year transformation program Bright Future is well on track



Significant changes in composition of leadership team

Cultural change addressed in three dedicated initiatives focused on customer centricity, market-driven innovation and corporate culture



Strategic roadmap materializing

---Measures for a bright future -----



Darmstadt

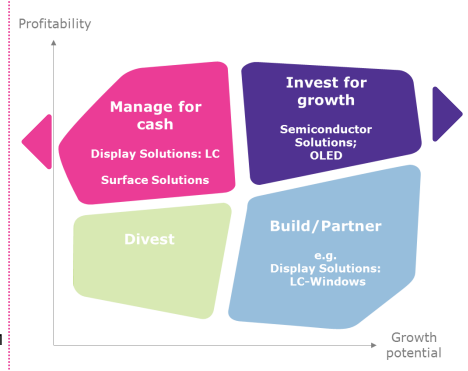
- In Darmstadt focus on R&D and production
- Immediate bottom line contribution from 2019 onwards
- Reduce the number of FTEs by ~15%
 = ~400 FTEs



 Chilworth site during September 2019 successfully closed



- Shut down of Electronics activities at Atsugi site started (to be completed during 2021)
- R&D and production activities in Atsugi transferred and consolidated in other PM locations in Asia
- Consolidation of site structure in Japan





- Leading supplier of high-purity process chemicals, gases and equipment serving semiconductor manufacturers
- Track record of accelerated growth and industry leading profitability
- Creating a leading electronic materials player with attractive long-term prospect

INTERMOLECULAR®

- Leading in advanced materials innovation
- Acquisition to strengthen semiconductor technology offering
- Application specific materials expertise with that perfectly complement Group's business and technology portfolio



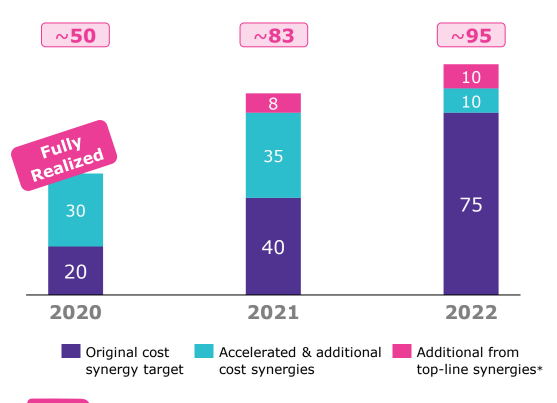
Both transactions successfully closed



Bottom-line management to support margin ambition of 30% in the long-term

Successful integration drives substantial synergy upgrade and acceleration

EBITDA pre impact of synergy ramp-up [€ m]



Sources of synergies



- Transform country setup
- Streamline duplicate structures



- Optimize production and supply chain network
- Achieve savings through joint procurement



- Integrate corporate & administrative functions
- Cost savings due to U.S. company delistings



Original target for 2022 is now being addressed for 2021

^{*}Top-line synergies from cross-selling, new products introductions and overarching initiatives

sustainability

06

Sustainability strategy enhanced, leveraging strengths with clear commitment to new targets

who we are

Targets

integration

Innovation Power

- We are a leading science and technology company with curious minds dedicated to human progress
- Long track-record in offering innovative products in attractive markets and serving important megatrends

Pioneering Products

- Well-equipped for developing new product classes: Portfolio of lifeimproving products in all businesses
- Enabling customers incl. scientists and developers to design next-gen products

Responsible Governance

- Resilient operations; sustainable leadership and risk-mitigation approach
- Responsibility is in our DNA: reflected by legal form, corporate governance and long history of more than 350 years

Goal #1: **Dedicated to human progress**

In 2030, we will advance human progress for more than 1 bn people through sustainable science & technology.

- and technology for our customers
- Sustainable innovations
 Impact of our technologies and products on health and well-being

Integrated

- Part of the overall strategy
- Linked to steering and operations
- Currently built into part of Executive Board compensation system

Goal #2: Creating sustainable value chains

By 2030, we will integrate sustainability into all our value chains.

- Sustainability
 Sustainable culture & values and transparent supply chain
- Securing our social license to operate in all regions

Steered & Reviewed

- Executive Board
- Supervisory Board
- Corporate Sustainability Committee

Goal #3: Reducing our ecological footprint

By 2040, we will achieve climate neutrality and reduce our resource consumption.

- Climate change & emissions
- Water & resource intensity

Communicated

- Development and reporting of meaningful KPIs
- Annual Report, Sustainability Report
- Investor events

Potential to increase sustainable value for business and society

and benefit High-IMPact SDGs where we can contribute Goal 3 GOOD HEALTH AND WELL-BEING **Business opportunities** Good Health > We are able to contribute with dedicated Develop a new range of sustainable products & services, and products, know-how, partnerships and benefiting from ourinnovation power initiatives in pharma, science and technology. Well-being Open up additional customer groups and expand regional reach Decent Work ➤ Our ambition of future growth considers and Economic health and safety of employees also in the **Risk management** Growth supply chain. Reduce risks through higher awareness and longer-term view Secure supply chain resilience Industry, Our innovation power will lead to more Innovation **Partnerships** sustainable products and processes in various and industries. 2 Contribute as supplier of choice to customers' ESG strategy Infrastructure Improve ESG impact of our suppliers RESPONSIBLE CONSUMPTION Responsible Being a responsible supplier, we will also 2 • Increase depth, meaning, and strategic focus of partnerships Consumption challenge suppliers to support in reaching and **Operations** 3 company targets. Production Increase attractiveness as employer 17 PARTNERSHIPS FOR THE GOALS Reduce costs of capital To unleash even more power, we foster **Partnerships** collaborations with capable partners to sum Benefit from grants and reliefs (politics, insurance, etc.) for the Goals 3 up know-how for more sustainable impact. Incentivize through integrated compensation schemes

Reduce our environmental footprint: Environmental targets 2020 have been achieved, new targets set

Achievements 2020

Reduce scope 1+2 emissions



2020 target¹: result¹: -20% -25%

Emissions target 2020 achieved!

25% overall reduction for Scope 1 and 2 emissions in 2020 relative to 2006 (planned: 20%)

Reduce water in stressed areas



Water target 2020 achieved!

- Water use in stressed areas reduced by 27% in 2020 vs. 2014 (planned: 10%)
- ✓ By 2020, all production sites⁴ successfully implemented sustainable water management system

Reduce Group Waste Score



2025 target³: result³: -5% -4.6%

Waste target ongoing & on track!

Based on Group Waste Score, reduced environmental impact by 4.6% vs. 2016 (planned: 5% by 2025)

 1 versus 2006 baseline, excluding Versum Materials

²versus 2014 baseline ³versus 2016 baseline

 4 Sites > 70.000 m 3 /a

New targets from 2021

- Aiming for climate neutrality (scope 1 to 3 emissions) by 2040
- Lower scope 1 and 2 GHG⁵
 emissions by 50% and to source
 80% of purchased electricity from
 renewable sources until 2030 vs.
 2020 baseline
- Absolute reduction of 1,500 kt⁶
 scope 3 CO₂ equivalents by 2030
- Enhancing water efficiency and improve the new Group water intensity score by 10% by 2025 vs. 2019 baseline
- Minimize negative environmental impacts, harmful emission residues should be lowered below a scientifically defined threshold by 2030

⁵GHG = Greenhouse Gas ⁶corresponds to ~30% of 2019 scope 3 emissions (current estimation incl. Versum Materials)



Next steps towards achieving ESG targets

AGENDA 2020-2022

Analysis of requirements: Strategy, business, regulation, stakeholders

Develop SBV tool² to measure product sustainability value

Link ESG¹ to board compensation

Build effective data platform for internal steering

Develop ESG KPIs for reporting

Further incorporate ESG in R&D, controlling, M&A and supply chain

Decide on dedicated investments and initiatives to achieve targets





Dedicated to human progress

Creating sustainable value chains

Reducing our ecological footprint

¹ESG: Environmental, Social, Governance

²Sustainable Business Value: Dive in deeper and read the research article on the <u>SBV method</u>



Executive Sumary

07

Latest COVID-19 assumptions for 2021

Overarching assumptions

- Increasing vaccination penetration across large populations in all major regions as of summer
- Global gradual easing of lockdowns to continue; vaccination efforts expected to keep up with virus mutations
- Overall improvement in the course of 2021 to continue; however, higher degree of forecast uncertainty



Healthcare assumptions

- Confirm ~ stable organic base business & pipeline sales target, despite higher uncertainty
- Pandemic impact on ramp-ups (particularly still depressed MS dynamic & high efficacy market in Q1 2021) remains a watch out; recent vaccination data expected to accelerate market position of Mavenclad
- Fertility to continue recovery

Life Science assumptions

- Continued strong additional demand & capacity expansions to support strong growth in Process Solutions
- Research and Applied more volatile and differentiated across customer and product segments; tailwinds for Research, about neutral effect in Applied

Electronics assumptions

- Neutral to positive impact on Semiconductor Solutions end markets
- Display and Surface Solutions to return to underlying trajectories



Full-year 2021 guidance

Net sales:

Organic: +10% to +12% YoY

FX: -2% to -4% YoY ~€18.5 - 19.5 bn

EBITDA pre:

Organic: +16% to +20% YoY (excl Biogen¹)

FX: -2% to -4% YoY ~€5.4 - 5.8 bn

EPS pre: ~€7.50 - 8.20



¹Q3 20 reversal of the provisions for the patent litigation proceedings for Rebif in the amount of ~€365 m; Guidance including Biogen – organic: +9% to +12%

Executive SUMMary



steady earnings growth with high wargins and a low risk profile







Successfully driving transformation into a leading science and technology company

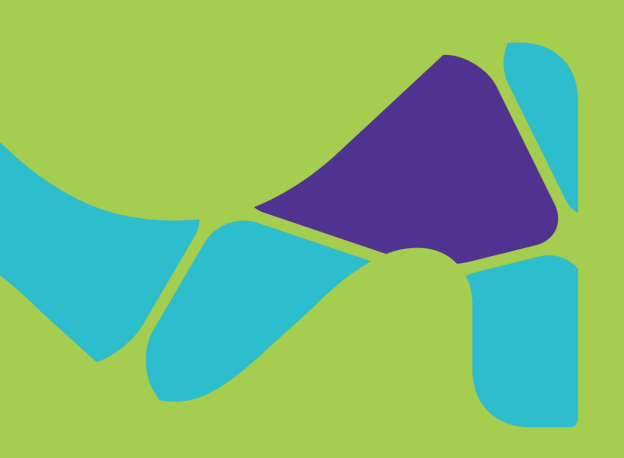
setup

Three-pillar structure strengthened further as a resilient basis; COVID-19 crisis as another proof point

Growth Engines

Healthcare pipeline, Process Solutions and Semiconductor Solutions will be key drivers of growth to 2022 and beyond Execution

Delivery on strategic priorities ensures profitable growth; regaining financial flexibility with higher likelihood of regular bolt-ons post 2022



xibneqqa



Successful crisis management increasingly mitigates pandemic impact



Guidance upgrade proof point of excellent crisis management and strong business performance

Group

2020 – strong resilience in times of global crisis

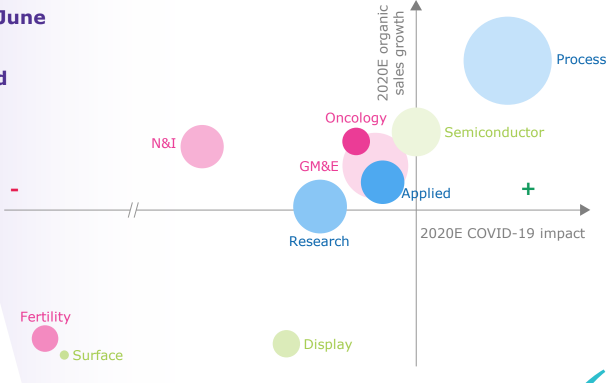


- Most businesses growing despite COVID-19
- Largest business growing and positively affected
- Smallest businesses with biggest impact

Delivery on priorities during crisis

- √ Health & safety of employees
- ✓ Business continuity
- ✓ Contributions to public health and society
- ✓ Sustainability aspects further enforced

Growth and COVID-19 impact by business¹





Merck KGaA, Darmstadt, Germany - steady earnings growth with high margins and a low risk profile



¹ Indicative only and based on guidance from August 6: slight to moderate organic sales and EBITDA pre growth, COVID-19 with up to a mid single-digit impact on sales of which 50-60% hitting EBITDA pre



2021 business sector guidance¹

Healthcare Life Science

Net sales

- Organic: +7% to +10%
- Mainly driven by Mavenclad[®], Bavencio[®] and recovery of Fertility
- Base business organically around stable

EBITDA pre

- Organic: +12% to +15% YoY (excl Biogen²)
- FX: -5% to -7% YoY
- ~€2,000 2,100 m

Net sales

- Organic: +15% to +18%
- Process Solutions as main growth driver

EBITDA pre

- Organic: +22% to +26% YoY
- FX: -1% to -3% YoY
- ~€2,850 3,000 m

Electronics

Net sales

- Organic: +5% to +7%
- Strong Semiconductor Solutions contribution, subject to quarterly DS&S project phasing
- OLED with high growth

EBITDA pre

- Organic: +9% to +12% YoY
- FX: -3% to -5% YoY
- ~€1,050 1,130 m



Additional financial guidance 2021

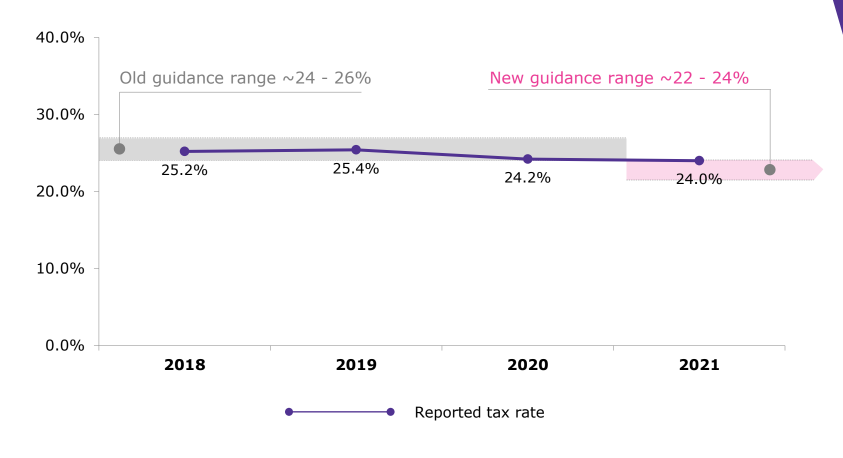
Further financial details

Corporate & Other EBITDA pre	~ €-440 to -490 m
Interest result	~ €-220 to -245 m
Effective tax rate	~22% to 24%
Capex on PPE	~€1.4 to 1.5 bn
Hedging/USD assumption	FY 2021 hedge ratio ~70% at EUR/USD ~1.17
2021 Ø EUR/USD assumption	~1.19 to 1.23



Underlying tax rate guidance lowered to new range of 22% to 24%

Tax rate development 2018-2020 and from 2021 onwards



Rationale for update

Strong profit growth in Life Science results in different profit contributions worldwide, leading to a lower overall tax rate



Group

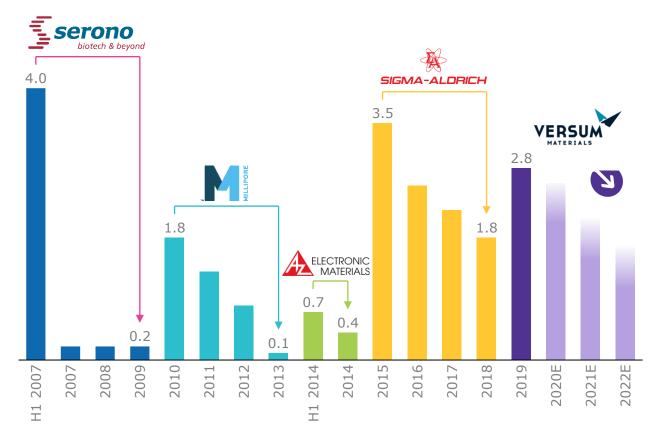
Focus on organic growth and deleveraging to 2022

Proven swift deleveraging after major acquisitions



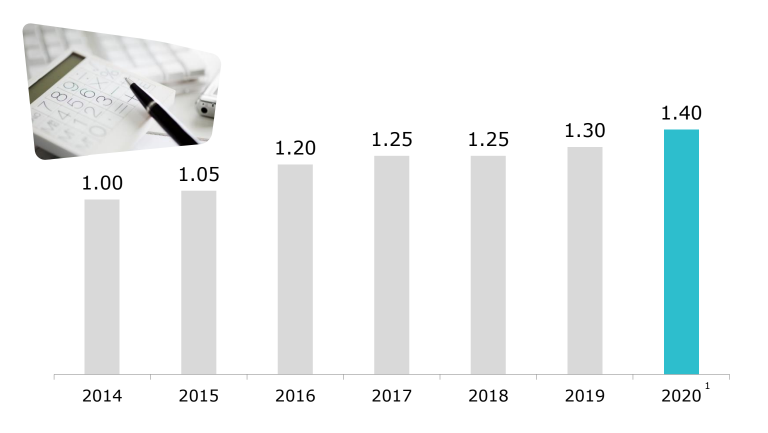
- Deleverage to <2x net debt/EBITDA pre in 2022
- M&A on hold until 2022; only smaller deals to be realized if budget available
- New mid-term capex ceiling of ~€1.3 bn reflects increased focus on organic investment and Versum consolidation
- Dividend policy mirrors
 sustainable earnings trend

Net debt / EBITDA pre track record & outlook



Sustainable dividend growth

Dividend¹ development 2014 -2020



2020 dividend

- Dividend of €1.40 (+8% YoY) per share proposed¹ for 2020
- Payout ratio of 23.1% of EPS pre² in 2020; aiming for 20-25% of EPS pre
- Dividend yield³ of 1.0%

³Calculated with 2020 year-end share price of € 140.35 per share.



¹Final decision is subject to Annual General Meeting approval

²Excluding Biogen provision release, including the provision release the ratio is 20.9%

Governance

Merck's KGaA, Darmstadt, Germany ownership structure

Shareholders of Merck KGaA, Darmstadt, Germany

- Shareholders hold ~30% of Group's total capital
- Share capital is divided into 129,242,252 bearer shares and one name share without nominal value
- Listed at German Stock Exchange's Prime Standard, member of the DAX 30
- Only bearer shares entitled to vote at Group's Annual General Meeting: one share, one vote

Supervisory Board

Its Chairman leads AGM, representing interests of Group's shareholders:

- Approving financial statements of the Group
- Working together with Executive Board, receiving reporting on progress regarding business and financial development at the Group, Containing 16 members:
 8 employee representatives, 8 independent representatives (2 family representatives and 6 elected shareholder representatives)
- 2 committees: Nomination and Audit Committee

Nomination Committee

Defines criteria and makes
proposals for new candidates

Audit Committee Control systems, external auditing, financial statements Finance Committee financial statements, external auditing

70%

Partners' Meeting & Family Board E. Merck KG, Darmstadt, Germany (99.9% Family) holds ~70% of Group's total capital

E. Merck KG, Darmstadt, Germany:

- These 70% of the total capital are not entitled to vote at Group's AGM
- A Family Board represents the entrepreneurial interests of the Family (elected by partners' meeting)

Board of Partners

Elected by Family Board, and takes over some duties of a usual Supervisory Board:

- Appointment, dismissal and supervision of Executive Board members of the Group
- Approves extraordinary business transactions and annual financial statements of E. Merck KG, Darmstadt, Germany
- Contains 5 members of the Family Board and 4 external members of the business community
- 3 committees: Finance, Personnel and R&D Committee

Personnel Committee
Executive board members,
contracts, compensation

R&D Committee research activities of all sectors: HC, LS, EL



Executive board compensation

Pay for performance reflecting the company's long-term strategy

		Long-Term Incentive Plan
ariable	40-50%	 Reflecting the long-term strategy for Group's growth and (from 2022) sustainability ambition 4 years performance cycle: 3 years target achievement + 1 year holding period Based on virtual Group Share Units (Grant € divided through start share price, multiplied with the end share price) Financial targets: 50% Company Share Price vs. DAX + 25% EBITDA pre margin + 25% Organic sales growth From 2022 multiplied with sustainability factor (0.8-1.2) reflecting KPIs from each of the sustainability goals Corridors for each target and achieved targets published transparently ex-post in the compensation report Maximum cap: Maximum pay out 250%, maximum € cap for LTIP for each board member published Claw-back allows to retain amounts allocated from the Long-Term Incentive Plan
Va		Profit Sharing Profit Sharing
	25-35%	 Three-years average profit after tax of the E. Merck Group KGaA, Darmstadt, Germany, multiplied with individual permille rate From 2021 reduced individual performance factor of 0.8-1.2 can increase (bonus) or decrease (malus) the amount based on a set of criteria, incl. the 3 sustainability goals, disclosure of catalogue and reasons for if performance factor ≠1.0 Individual permille rate for each board member and maximum € cap for each board member published Staggered incentivization and minimum threshold value and maximum limit for profit after tax (0.75/2.0 bn €)
_	6-9%	 Mandatory personal investment in Company Shares amounting to one third of the net payment of the profit sharing (4 year holding period) Pension Entitlements Defined contribution
	0-3%	Additional Benefits Mainly contributions to insurance policies, personal security expenses, company car
sic		Basic Compensation
Ba	15-20%	 Fixed and non-performance related compensation Paid in 12 equal monthly installments 1.4 million € for the chairman / up to 1.1 million € for the members of the executive board
_	Maximum	total compensation: reduced to €11.5 m Chairman, €9.5 m other executive board members

External stakeholders assess our engagement



As of 2020, Merck KGaA, Darmstadt, Germany received an MSCI ESG* Rating of AAA.

*Environment, Social, Governance



2021, we received an **ESG** Risk Rating of 19.5 and Sustainalytics: low risk of experiencing material financial impacts from ESG factors.

PLATINUM

ecovadis



annually examines ~75,000



measuring the performance

FTSE4Good

Since 2008, Company is part

of FTSE4Good Index,

access to medicine = Index **T**

In the 2021 Access to **Medicine Index** Group ranked eighth place. We were recognized for our performance in R&D, where we ranked fifth.



In 2020, Merck KGaA, Darmstadt, Germany has once more achieved prime status by ISS Oekom.

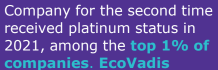


In 2019, the Group share was again included in **STOXX Global ESG Leaders** Index.

a sustainability index based on key environmental, social and governance criteria.



Company has been reconfirmed as a constituent of the **Ethibel Sustainability Index (ESI) Excellence Europe** since May 2020, based on VigeoEiris.



suppliers from 160 countries.



CDP Climate: In 2020, we scored "B" (2019: C). CDP Water: In 2020, we received a "B" (2019: B).

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Group

Regular portfolio review remains key to success

strong track record

- Acquisitions and divestments are part of Group's history
- Licensing and partnerships remain on our agenda
- All prior transactions earned their cost of capital



pefining portfolio guard rails

- Three strong pillars with no business marginalized
- Leading market position in attractive markets
- Focus on innovation and sustainability through science and technology



clear financial M&A criteria

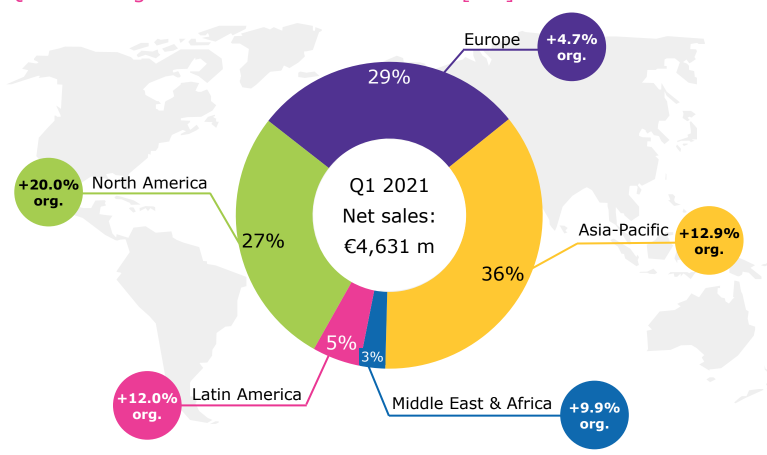
- Supporting profitable growth strategy
- IRR > WACC
- EPS pre accretive
- Maintain investment grade rating



- Current set-up is strong and organic investment opportunities are attractive
- Expect to regain financial flexibility by 2022 to pursue external growth opportunities
- Targeted and more regular bolt-on approach more likely than large transformative deals

Life Science demand drives particularly strong growth in North America and Asia-Pacific

Q1 2020 Regional breakdown of net sales [€m]



Regional organic development

- APAC: strong Life Science and Healthcare growth while Electronics
 stable with Semi offsetting Display
- Europe: Growth in Process & Research Solutions more than compensates Healthcare decline largely in CM&E
- North America: growth across all three sectors, particularly strong Life
 Science and Bavencio[®] performance
- LATAM growth driven foremost by Fertility and Bavencio[®]
- CM&E is the largest growth driver in ME&A



27% organic growth in Life Science, moderate Healthcare growth & stable Electronics drive very strong organic Group sales and EBITDA pre growth

Q1 YoY Net Sales	Organic	Currency	Portfolio	Total
Healthcare	3.5%	-5.9%	-1.2%	-3.6%
Life Science	26.7%	-6.2%	0.0%	20.4%
Electronics	0.2%	-4.5%	0.0%	-4.3%
Group	12.2%	-5.8%	-0.4%	6.0%

- 22% org. growth in Fertility and 20% org. growth in Oncology more than compensate for N&I decline (-4% org.) amid depressed dynamic market and VBP impact in CM&E¹ (-4% org.)
- Record Life Science organic growth driven by all business units with Process Solutions up +38%; Research elevated to +24%; Applied Solutions delivers 8% organic growth
- Electronics about stable as Semiconductor Solutions growth (+4% org.) and recovering Surface Solutions (+5% org.) are offsetting decline in Display (-7% org.)

Organic EBITDA pre growth three times faster than sales growth

Currency

Portfolio

Q1 2020

Organic

- Strong Life Science gross profit further boosted by Bavencio[®] milestones and continued cost discipline in all sectors
- FX burden of -8% across various currencies with largest negative impact from USD, BRL and JPY; partly mitigated by hedging

Q1 2021

Q1 YoY EBITDA pre

36.3% €1,511 m

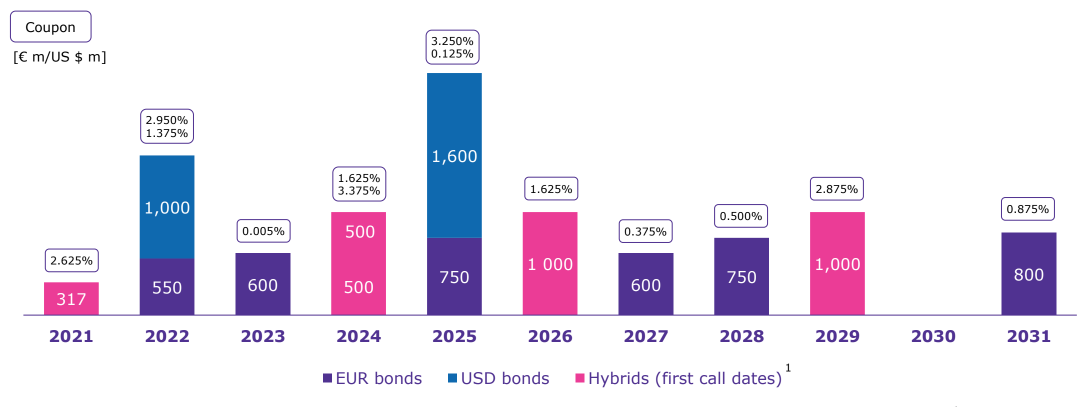
€1,181 m -8.3% -0.1%

M

¹ Cardiovascular, Metabolism and Endocrinology (new Franchise name as of Q1 2021)

Balanced maturity profile: Lower refinancing risks & higher flexibility

Maturity profile as of March 31, 2021



¹No decision on call rights taken yet



Q1 2021: Overview

Key figures

[€m]	Q1 2020	Q1 2021	Δ
Net sales	4,370	4,631	6.0%
EBITDA pre Margin (in % of net sales)	1,181 <i>27.0%</i>	1,511 32.6%	27.9%
EPS pre	1.50	2.18	45.3%
Operating cash flow	516	1,216	135.4%
[€m]	Dec. 31, 2020	March 31, 2021	Δ
Net financial debt	10,758	10,081	-6.3%
Working capital	3,938	4,231	7.4%
Employees	58,096	57,933	0.0%

Comments

- Net sales growth of 6% driven by 12% organic growth and FX burden of -6%
- EBITDA pre increase, driven particularly by operating leverage in Life Science further boosted by Bavencio[®] milestones
- EPS pre driven by EBIT pre growth, supported by better financial result & lower effective tax rate
- Operating cash flow more than doubles, largely driven by strong EBITDA pre growth and favorable net working capital
- Significant reduction of net financial debt



Q1 2021: Reported figures

Reported results

[€m]	Q1 2020	Q1 2021	Δ
EBIT	716	1,043	45.7%
Financial result	-98	-59	-40.0%
Profit before tax	617	984	59.4%
Income tax	-159	-236	48.1%
Effective tax rate	25.8%	24.0%	
Net income	456	747	63.7%
EPS (€)	1.05	1.72	63.8%

Comments

- Strong performance across all sectors particularly in Life Science drives 46% EBIT growth
- Reduced interest expense and lower LTIP provisions drive improved financial result
- Effective tax rate at the higher end of the new guidance range
- EBIT growth, improved financial result and lower tax rate drive higher net income & EPS



Cash flow statement

Q1 2021 – cash flow statement

[€m]	Q1 2020	Q1 2021	Δ
Profit after tax	458	748	290
D&A	431	424	-7
Changes in provisions	16	-34	-50
Changes in other assets/liabilities	-23	160	183
Other operating activities	-10	6	16
Changes in working capital	-356	-88	267
Operating cash flow	516	1,216	700
Investing cash flow	-288	-346	-58
thereof Capex on PPE	-337	-309	29
Financing cash flow	542	6	-536

Cash flow drivers

- Strong increase in profit after tax driven particularly by Life Science, further boosted by Bavencio[®] milestones
- Provisions largely reflect various favorable developments in litigation positions
- Contribution from other assets/liabilities largely explained by tax positions
- Working capital upside mainly driven by higher payables (Q4 2020 phasing effect)
- Delta in investing cash flow primarily explained by divestment of Allergopharma in Q1 2020
- Capex driven foremost by Life Science capacity expansions



Adjustments in Q1 2021

EBIT Adjustments

[€m]	Q1 20	020	Q1 20)21
	Adjustments	thereof D&A	Adjustments	thereof D&A
Healthcare	-27	2	10	0
Life Science	11	0	14	0
Electronics	35	0	17	3
Corporate & Other	17	0	6	0
Total	36	2	47	3





Financial calendar

Date	Event
May 12, 2021	Q1 2021 Earnings release
August 5, 2021	Q2 2021 Earnings release
September 9, 2021	Virtual Capital Markets Day
November 11, 2021	Q3 2021 Earnings release
April 22, 2022	Annual General Meeting



нealthcare

China's VBP: Round 4 bidding recently completed, sustained confidence in keeping base business approx. stable through 2021 and 2022



Round 1		Round 2	Rou	nd 3	Round 4
2019			2020		DATE 2021
March: First round initiated, no Merck's KGaA, Darmstadt, Germany products impacted	December: Round 2 announced (incl. bisoprolol)	April: Round 2 winners granted exclusive access to 60% of total hospital market ¹ , non- winners (incl. Concor®) retain access to remaining 40% of hospital market at a gradient price cut ² + can freely compete in non-hospital/ retail market	July: Round 3 announced (incl. Metformin IR and XR) ¹	August: Bidding manufacturers), (• Access retained market (~30% of the hospita) • Continued grown	Jan: Round 4 announced (incl. calcium dobesilate³ and canagliflozin³), bidding completed on Feb 3, Invokana® & Doxium ® not among winners, Implementation starts from late April/May subsequentially completed, 8 winners (all Chinese Glucophage® not among winners ed to non-hospital/retail of total market), and ~20% I market (at gradient price cut) with of Chinese metformin p.a. over the next 5 years



China Glucophage sales represent only ~8% of the total base business (2020 net sales)
Sustained confidence in approx. stable base business (org.) through 2021 and 2022

^{1:} hospital market for bisoprolol and metformin makes up ~70% of total market, this includes urban hospitals, rural hospitals, and community health centers; 2: Concor® price cut in the high single digit %; 3: alliance products; Acronyms: VBP = Volume-Based Procurement



Phase I

M1231 Bispec. MUC1xEGFR ADC Solid tumors

M1774 ATR inhibitor Solid tumors

peposertib DNA-PK inhibitor Solid tumors¹ bintrafusp alfa
TGFbeta trap/anti-PD-L1
Cervical cancer 1L

M6223 anti-TIGIT mAb Solid tumors²

M5049 TLR7/8 antagonist Systemic lupus erythematosus / Cutaneous lupus erythematosus

M5717 PeEF2 inhibitor

Simplified overview excluding programs for which Merck KGaA, Darmstadt, Germany explores externalization opportunities and/or for which Company pursues only limited internal development activities

Phase II

berzosertib ATR inhibitorSmall-Cell Lung Cancer³

tepotinib
MET kinase inhibitor
Metastatic Colorectal Cancer
RAS/BRAF wt, MET amplified⁴

tepotinib
MET kinase inhibitor
Non-small cell lung cancer,
EGFR mutant, MET amplified⁵

bintrafusp alfa TGFbeta trap/anti-PD-L1 Non-small cell lung cancer 1L/2L

bintrafusp alfa TGFbeta trap/anti-PD-L1 Locally advanced non-small cell lung cancer bintrafusp alfa TGFbeta trap/anti-PD-L1 Biliary tract cancer 1L

bintrafusp alfa
TGFbeta trap/anti-PD-L1
Cervical cancer 2L

bintrafusp alfa TGFbeta trap/anti-PD-L1 Triple negative breast cancer (HMGA2 positive)

M5049 TLR7/8 antagonist COVID-19 pneumonia

Phase III

xevinapant IAP inhibitor

Locally advanced squamous cell carcinoma of the head and neck⁶

avelumab anti-PD-L1 mAb Non-small cell lung cancer 1L

evobrutinib BTK inhibitorRelapsing multiple sclerosis

Registration

tepotinib MET kinase inhibitor Non-small cell lung cancer, METex14 skipping^{7,8}

avelumab anti-PD-L1 mAb Urothelial cancer 1L-M^{9,10}

ADC: Antibody Drug Conjugate; Bispec.: bispecific; It. first-line treatment; 1L-M: first-line maintenance treatment; 2L: second-line treatment;

¹ Study in combination with avelumab. ² Includes study in combination with bintrafusp alfa. ³ Includes studies (phase I/II) in collaboration with/ sponsored by external partners, e.g. NCI. ⁴ In combination with cetuximab. ⁵ In combination with osimertinib. ⁶ On March 01, Merck KGaA, Darmstadt, Germany announced a worldwide in-licensing agreement with Debiopharm, Switzerland, for the development and commercialization of xevinapant (Debio 1143). ⁻ As announced on February 03, 2021, the US Food and Drug Administration (FDA) has approved tepotinib for the treatment of adult patients with metastatic non-small cell lung cancer (NSCLC) harboring mesenchymal-epithelial transition (MET) exon 14 skipping alterations. This indication is approved under accelerated approval based on overall response rate and duration of response. Continued approval for this indication may be contingent upon verification and description of clinical benefit in confirmatory trials. ⁸ As announced on November 26, 2020, the European Medicines Agency (EMA) has validated for review the application for tepotinib for the treatment of adult patients with advanced non-small cell lung cancer. ⁹ As announced on January 25, 2021, the European Commission (EC) has approved avelumab as monotherapy for the first-line maintenance treatment of adult patients with locally advanced on metastatic urothelial carcinoma. ¹⁰ As announced on February 24, 2021, Japan's Ministry of Health, Labor and Welfare (MHLW) has approved a new indication for avelumab as a first-line maintenance treatment for advanced bladder cancer. Additional information: Several combination studies (phase II) of avelumab with talazoparib, axitinib, ALK inhibitors or chemotherapy ongoing under sponsorship of Pfizer.

Unless noted otherwise, clinical programs conducted in collaboration with external partners are not shown unless Merck KGaA, Darmstadt, Germany has co-ownership of data. In such case the indication is shown in *Italics*.

Pipeline products are under clinical investigation and have not been proven to be safe and effective. There is no guarantee any product will be approved in the sought-after indication.

Oncology

Immuno-Oncology

Immunology

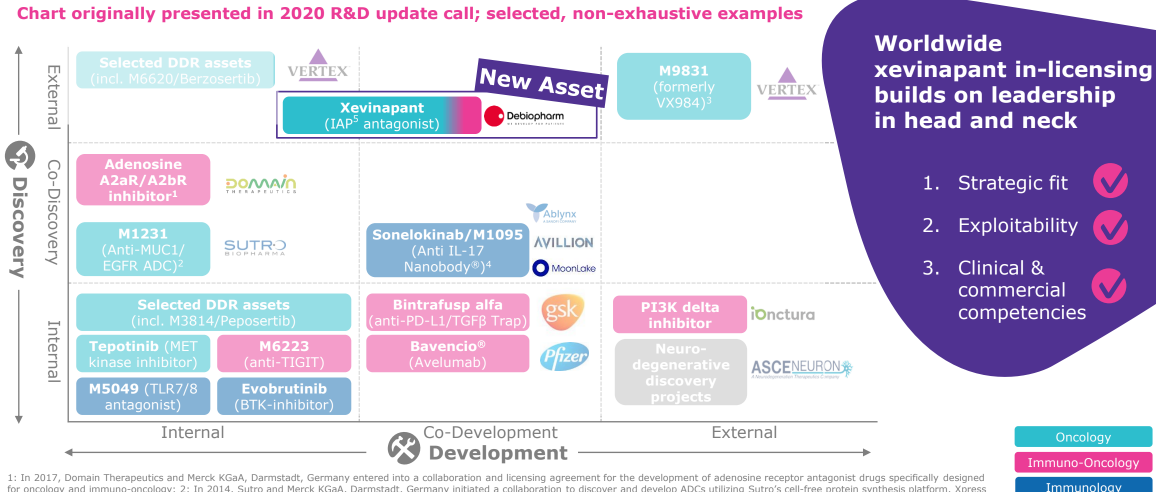
Neurology

Global Health



Xevinapant

Potential to become standard of care in core area for the Group



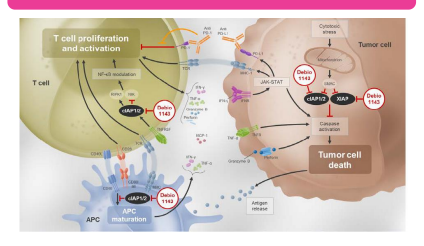
1: In 2017, Domain Therapeutics and Merck KGaA, Darmstadt, Germany entered into a collaboration and licensing agreement for the development of adenosine receptor antagonist drugs specifically designed for oncology and immuno-oncology; 2: In 2014, Sutro and Merck KGaA, Darmstadt, Germany initiated a collaboration to discover and develop ADCs utilizing Sutro's cell-free protein synthesis platform, Xpress CF+[™]. Merck KGaA, Darmstadt, Germany is responsible for drug product, clinical development and commercialization of any resulting products; 3: In 2019, an exclusive license was granted to Vertex for the use of M9831 in gene-editing applications; 4: Avillion conducted Ph II of M1095 in Psoriasis, Merck KGaA, Darmstadt, Germany decided to out license sonelokinab to a new partner to initiate Phase III development in 2021 5: Inhibitor of Apoptosis Proteins



Xevinapant (Debio 1143)

Potentially first in class oral IAP antagonist with FDA BTD

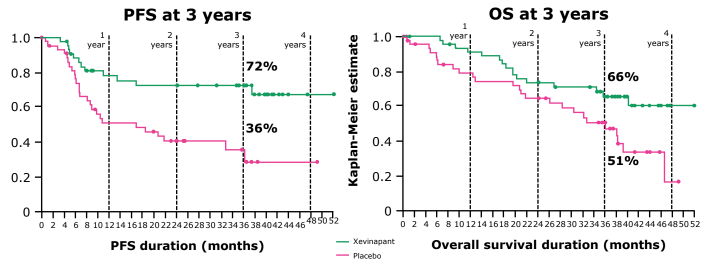
Mode of Action¹



- Oral Inhibitor of Apoptosis Proteins (IAP) antagonist: chemo-/radio-sensitizer & enhancer of anti-tumor immunity
- IAP antagonists tackling two cancer hallmarks:
 - Enhancing anti-tumor immunity
 - Lowering threshold for tumor cell death

Compelling Phase 2 data² published in *The Lancet Oncology*, and presented at ESMO 2020

- Improvement in OS statistically significant and clinically meaningful: HR 0.49 (0.26-0.92); p=0.0261
- Clinically compelling PFS improvement: HR 0.34 (0.17-0.68); p=0.0023
- Predictable and manageable safety profile without substantial additional toxicity to standard CRT



Acronyms: BTD = Breakthrough Therapy Designation; IAP = Inhibitor of Apoptosis Proteins; 1: Debiopharm; 2: ESMO 2020 - Late Breaking Abstract 39 - 3-years follow-up of double-blind randomized phase II comparing concurrent high-dose cisplatin chemo-radiation plus xevinapant or placebo in high-risk patients with locally advanced squamous cell carcinoma of the head and neck



Xevinapant

Total deal-volume of up to ~ €900 m and industry-typical sales royalties

Payment type	Amount (in €)	Accounting treatment ²	
Upfront payment	~ €190 m	Largest part to be capitalized as an intangible asset	
Approval milestones	Up to ~ €380 m ¹	To be paid and capitalized as an intangible asset upon approval and to be amortized once asset is ready for use	
Commercial milestones	Up to ~ €330 m	To be paid and capitalized as an intangible asset, based on sales thresholds and to be amortized over remaining useful life	
Sales	n/a	Group to recognize sales globally (incl. US)	
R&D Costs	n/a	For ongoing TrilynX study Cash view: 50/50 cost sharing P&L view: fully shown in Group P&L 2nd study for cisplatin-ineligible patients: Company incurs 100% of cost	
Royalties	n/a	Group to pay industry-typical sales royalty to Debiopharm	

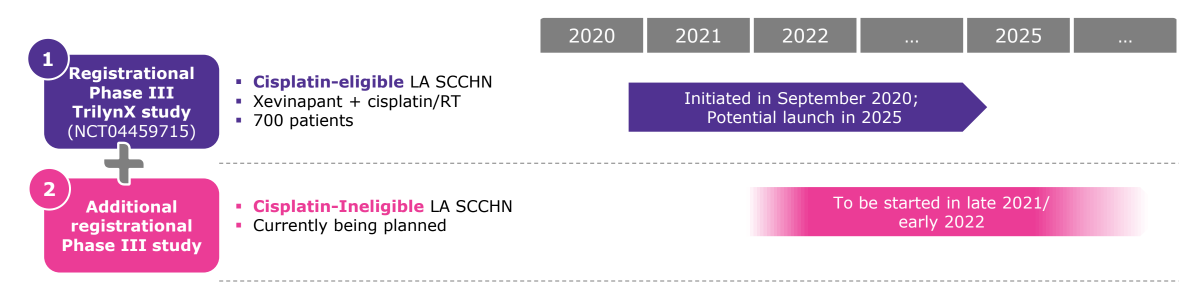
 $^{^{1}}$ thereof up to \sim €300 m for focus H&N indications)



 $^{^{\}rm 2}$ final accounting treatment is still subject to alignment with auditors

Xevinapant (Potentially first in class oral IAP antagonist)

Two Phase III studies are designed to target the majority of unresectable LA SCCHN patients receiving systemic therapy + RT





Blockbuster potential provided success of both studies



20,000+ unresectable LA SCCHN patients in US and EU-5 each



Tepotinib (MET kinase inhibitor)

First-in-class launch in MET Exon14 sets foundation for EGFRm/ METamp opportunity and exploration in other tumor types





NSCLC - MET Exon14

(VISION study, NCT02864992 3-5% of total NSCLC population)

- Highly competitive data set presented at ASCO 2020 and published in New England Journal of Medicine in May 2020 (99 patients with a follow up ≥9 mths)
- First-in-class launch¹ in Japan² in March 2020, Sakigake designation³ granted in 2018
- Approval by US FDA¹ received on February 3, 2021



Tap into a growing opportunity in NSCLC - EGFRmut/ METamp

(INSIGHT 2 study, NCT03940703 2-5% of total NSCLC population)

- Increased EGFRm detection with testing and treatment moving into earlier lines of therapy (ADAURA trial demonstrates a 79% reduction in the risk of death with Osimertinib in the adjuvant setting (ASCO 2020), suggesting an even greater uptake of Osimertinib)
- **METamp as the primary driver of resistance** Some publications suggest that METamp resistance post-Osimertinib could be ~25%⁴



EXPLOTE EGFR resistance in crc – Tepotinib + Erbitux® combo(NCT04515394)

Opportunity for Tepotinib to address an unmet need in metastatic colorectal cancer (mCRC) together with Erbitux® ERBITUX

1: approved for both treatment naïve and previously treated METex14 positive NSCLC patients; 2: second largest Oncology market globally; 3: SAKIGAKE designation promotes research and development in Japan, aiming at early practical application for innovative pharmaceutical products; 4: Piotrowska et al., "Landscape of Acquired Resistance to Osimertinib in EGFR -Mutant NSCLC and Clinical Validation of Combined EGFR and RET Inhibition with Osimertinib and BLU-667 for Acquired RET Fusion", AACR Cancer Discovery 2018; Acronyms: CRC = Colorectal cancer; EGFR = Epidermal Growth Factor Receptor; NSCLC = Non-small cell lung cancer



Tepotinib (MET kinase inhibitor)

Tapping into the rapidly evolving EGFRmut/METamp market – Encouraging INSIGHT 1 data





INSIGHT 2 – Tepotinib + Osimertinib in Osimertinib Relapsed METamp NSCLC

- Study design recently amended to reflect evolved and future standard of care:
 - Target population Inclusion criteria adjusted to focus solely on 1L Osimertinib failures
 - Testing Streamline patient enrollment based on current gold standard method (TBx FISH)
 - Increasing METamp prevalence Some publications suggest that METamp resistance post-Osimertinib could be ~25%¹
- Estimated primary completion date: November 2022

Tepotinib + Erbitux® (Cetuximab) - Adressing a significant medical need in 2L metastatic colorectal cancer (mCRC)

- Opportunity for Tepotinib to address an unmet need in CRC together with Erbitux®
- Estimated primary completion date: March 2023

A solid foundation - Encouraging INSIGHT 1 data (18-months follow-up presented at WCLC 2019)²

Endpoint	Tepotinib + gefitinib	Chemotherapy
Primary - PFS (HR 0.13 [90% CI 0.04, 0.43])	16.6 m	4.2 m
Secondary - ORR (OR 2.67 [90% CI 0.37, 19.56])	66.7%	42.9%
Secondary - OS (HR 0.09 [CI 0.01, 0.54])	37.3 m	13.1 m

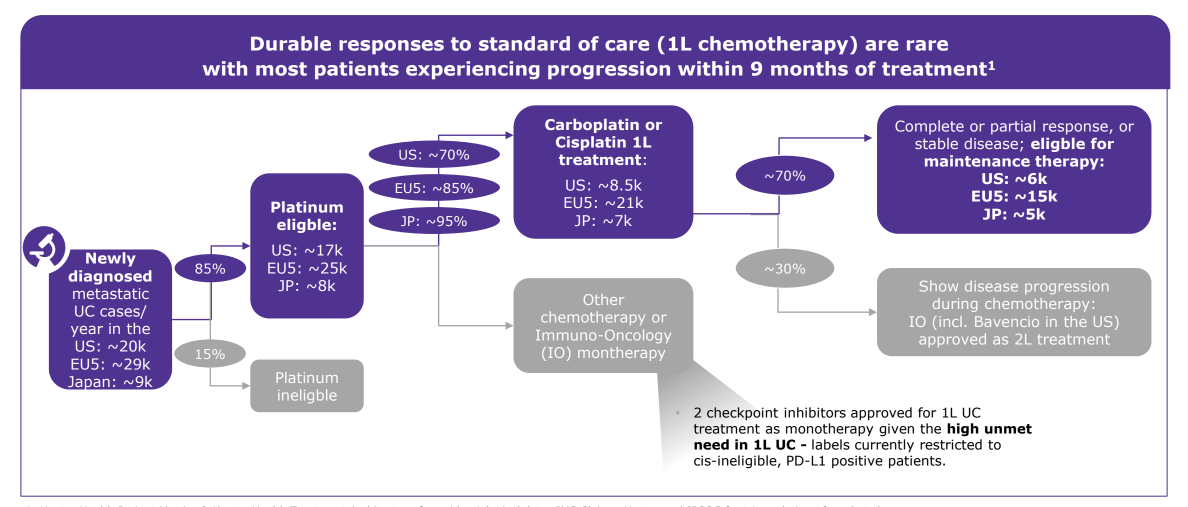


Safety: generally well-tolerated, most adverse events mild to moderate



1: Piotrowska et al., "Landscape of Acquired Resistance to Osimertinib in EGFR -Mutant NSCLC and Clinical Validation of Combined EGFR and RET Inhibition with Osimertinib and BLU-667 for Acquired RET Fusion", AACR Cancer Discovery 2018; 2: Wu et al., "Long term outcomes to tepotinib plus gefitinib in patients with EGFR mutant NSCLC and MET dysregulation: 18 month follow up", presented at WCLC 2019; Acronyms: FISH = Fluorescence in situ hybridization; TBx = Tissue Biopsy

Bavencio® (Avelumab) – Urothelial Carcinoma (UC 1L) UC 1L maintenance treatment achieving transformative OS benefit (31% reduction in risk of death, 7 months increase in median overall survival)





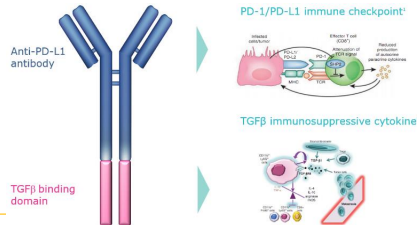


Bintrafusp alfa¹

An innovative first-in-class bifunctional fusion protein discovered in-house leading the TGF- β immuno-oncology field



- Innovative **first-in-class bifunctional fusion protein** designed to simultaneously target two immune suppressive pathways (blocking PD-L1 and reducing TGF-β signaling)
- Demonstrated superior anti-tumor activity in pre-clinical study compared to anti-PD-L1 alone, and anti-PD-L1 and TGF-β given in combination as separate agents
- Great excitement in IO community about M7824 uniquely addressing TGF-ß biology widely accepted as key resistance factor for anti-PDx therapies





- Tested in **14 Phase Ib expansion cohorts** across >700 patients in more than 10 tumor types
- Shown clinical anti-tumor activity across multiple hard-to-treat cancers including advanced NSCLC, biliary tract cancer, HPV-associated cancers, and gastric cancer
- PhII study M7824 monotherapy versus pembrolizumab 1L, advanced NSCLC high PD-L1-tumor expressers started in October 2018



 Multiple high priority immuno-oncology clinical development studies ongoing or expected to commence shortly, including studies in non-small cell lung and biliary tract cancers with registrational intent and most recently advanced, unresectable cervical cancer

Acronyms: NSCLC = Non-small Cell Lung Cancer, IO = Immuno-Oncology

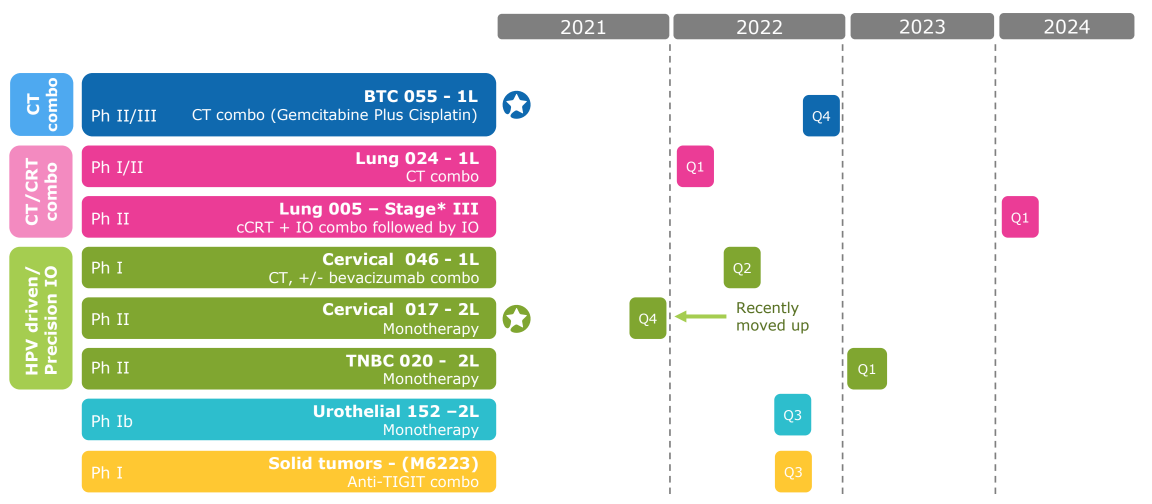


proposed International Nonproprietary Name (INN)

Bintrafusp alfa

INTR@PID Program: Upcoming Readouts





Acronyms: BTC = Biliary Tract Cancer; CT = Chemotherapy; EMT = Epithelial-mesenchymal transition; HPV = Human papillomavirus; NSCLC = Non-small Cell Lung Cancer; RT = Radiation therapy; TNBC = Triple-Negative Breast Cancer; * unresectable; All clinical timelines are event-driven and may be subject to change



Bintrafusp alfa: Developmental Progress

NSCLC Stage III cCRT Combo trial

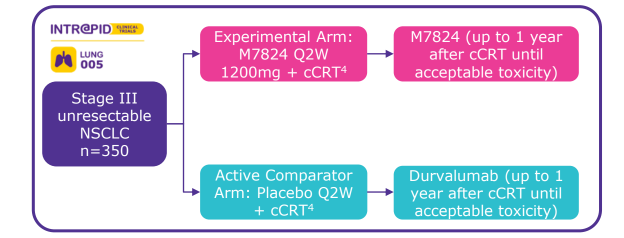
NSCLC 2L data presented at ESMO 2018

- Need: NSCLC accounts for 80-85% of all cases of lung cancer¹
- Results: Encouraging efficacy comparing favorably to established PDx-inhibitor monotherapy (IRC)2,3:
 - **ORR (all-comers):** 25.0%
 - ORR (PD-L1-positive): 37.0%
 - **ORR (PD-L1-high):** 85.7%
- Progression free survival by IRC (PD-L1 \geq 1%):
 - M7824: **mPFS = 9.5 months**, competitor: $4.0 \text{ months}^{2,3}$
- Overall Survival by IRC (PD-L1 \geq 1%):
 - M7824: mOS not reached, competitor: 12.7 months^{2,3}

Pre-clinical data on M7824 + RT combo⁵

- M7824 and RT combination therapy enhances antitumor activity relative to mono-therapies in mouse models
- EMT, VEGF, and RT-induced fibrosis gene signatures are decreased with M7824 and combination therapy, and M7824 reduces RT-induced fibrosis
- Results support evaluation of M7824 + RT in the clinic

INTR@PID LUNG 005



Endpoints

Primary endpoint: PFS

Main secondary endpoints: OS, Safety, Pulmonary function, Association of PD-L1 expression at base line and efficacy



Bintrafusp alfa: Developmental Progress

Data shown at AACR 2019 highlights opportunity in HPV-related cancers

Efficacy variable	HPV-associated cancer (n=43)	HPV+* (n=36)		
Confirmed BOR, n (%)				
CR	2 (4.7%)	2 (5.6%)		
PR	10 (23.3%)	9 (25%)		
SD	6 (14.0%)	5 (13.9%)		
PD	20 (46.5%)	17 (47.2%)		
Not evaluable	5 (11.6%)	3 (8.3%)		
Delayed PR ⁺	3 (7.0%)	3 (8.3%)		
ORR per RECIST v1.1, n (%) [95% CI]	12 (27.9%) [15.3–43.7]	11 (30.6%) [16.3-48.1]		
Total clinical response rate [†] , n (%)	15 (34.9%)	14 (38.9%)		
DCR, n (%)	18 (41.9%)	44.4%		

Prevalence: >630,000 new cases of HPV-related cancer are reported worldwide annually¹

Response Rates:

- Bintrasfusp alfa response rates compared favorably to those with anti-PD-1 inhibitors (ORRs of 13%-24%)¹⁻⁷
- ORR was 27.9% and 30.6% in HPV-associated and HPV+ cancers, respectively
- Including three additional patients with delayed PRs after initial PD: Total response rate was 34.9% and 38.9% in HPVassociated and HPV+ cancers, respectively

Long-term Benefit:

- Most responses durable with 4 responses having DoR >18 months and 11/15 responses ongoing at the data cutoff
- Responses to bintrafusp alfa occurred irrespective of tumor type or PD-L1 expression
- Safety profile was similar to anti-PD-(L)1 therapy^{1,5} except for SCC/KAs and low grade mucosal bleeding which are anticipated AEs with TGF-β inhibition^{8,9}

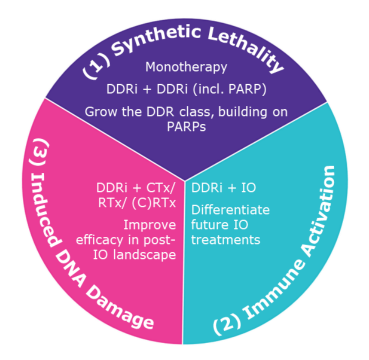
Cervical Cancer 2L study recently posted on ct.gov

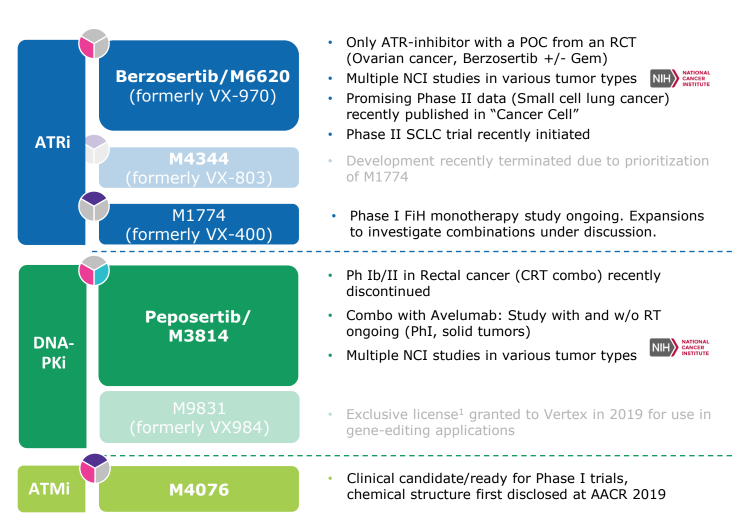


DNA Damage Response (DDR)

Leading DDR portfolio with a broad clinical program

Strategy presented at R&D Update Call 2019





1: incl. upfront payment + milestone/royalties on future sales; Acronyms: ATMi = Ataxia telangiectasia-mutated; ATRi = Ataxia telangiectasia and Rad3-related inhibitors; CRT = Chemoradiotherapy; DDR = DNA Damage Response; DNA-PKi = DNA-dependent Protein Kinase Inhibitor; PARP = poly(ADP-ribose) polymerase inhibitor; POC = Proof of concept; RCT = Randomized Controlled Trial; RT = Radiation Therapy

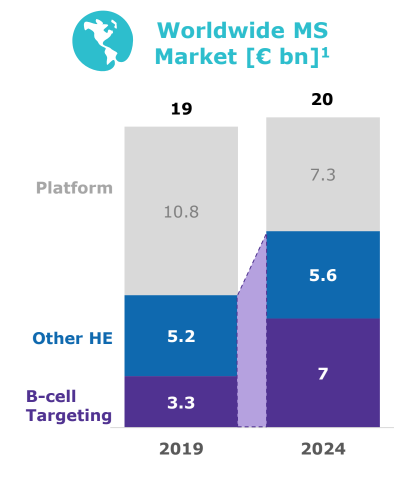


We pioneered BTKi development for MS with Evobrutinib

Potential to have 3 complementary MS branded products by 2025

Unmet need in Multiple Sclerosis (MS) – Need for new mechanisms to control disease

- ~50% of patients with Relapsing MS (RMS) continue to have ongoing disease activity over 2 years even when treated with the most effective agents
- No therapy with impact on progression mediated by CNS myeloid cells
- Systemic side effects of therapies limit patient acceptance and compliance
- All approved higher efficacy therapies associated with elevated risk of infection



B-Cell Targeting +
High-Efficacy (HE) Orals
represent >60% of MS sales

BTKi is a novel class of non-

depleting therapies selectively targeting both B-cells and innate immune cells including disease progression-relevant microglia

Merck KGaA, Darmstadt, Germany was the first to conduct a full

Phase II dose-ranging study in MS with Evobrutinib, a highly selective covalent BTKi²

Merck KGaA, Darmstadt,
Germany is a growing MS player
and could have 3 complementary
branded products by 2025 –
Mavenclad®, Rebif®, Evobrutinib

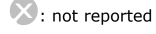
Platform agents – interferons, copaxone, DMFs and Teriflunomide; Other HE (high-efficacy) - cladribine, S1Ps, alemtuzumab; B-cell Targeting – ocrelizumab, ofatumumab, ublituximab. Includes branded products, generics and biosimilars; 1: Merck KGaA, Darmstadt, Germany internal estimates; 2: Montalban et al. NEJM 2019; 380:2406-2417; Acronyms: BTKi = Bruton's tyrosine kinase inhibitor

Evobrutinib stands out amongst BTK inhibitors under development Uniquely positioned both in terms of clinical evidence and mode of action

		Fenebrutinib##	Tolebrutinib**	Evobrutinib
0	Long-term* efficacy on relapses			(1)
Evidence	Long-term* safety			(1)
	Convenience (oral)	⊘ BID	₹ QD	▼ BID
Clinical	Exposure in CSF		(2, ##) in HV	(3) in MS
	Biomarker of inflammation and progression in MS patients (sNfL)			(3)
ical	BTK occupancy in the CNS		(4)	(5)
Preclinical data	Efficacy in progressive EAE model and reduction of leptomeningeal inflammation#			(6-8)

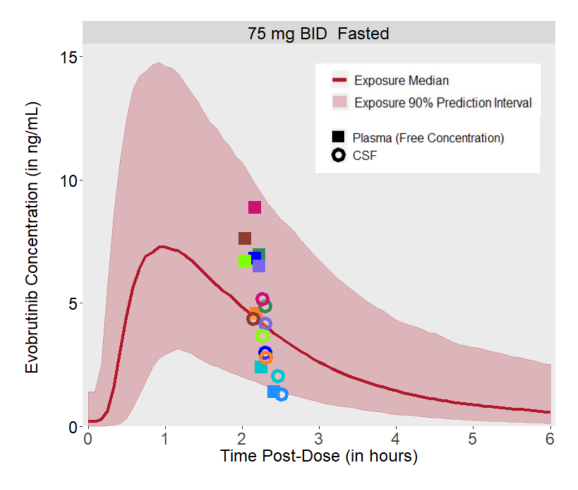


Phase III studies: Recruitment on track → Target data in-house in Q4 2023 and potential filing shortly after





Evobrutinib has the potential to tackle Multiple Sclerosis directly in the brain



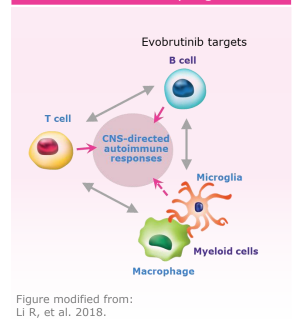
- Evobrutinib was quantifiable in CSF of 9 of 9 clinically stable RMS patients administered the efficacious 75 mg BID dose in the RMS Phase 2 OLE study*
 - Plasma PK were at steady-state and consistent with other RMS Phase2 patients
- CSF concentrations are consistent with free plasma concentrations
- The measured CSF concentrations are biologically relevant
 - Median BTK trough occupancy >95% observed at similar plasma concentrations



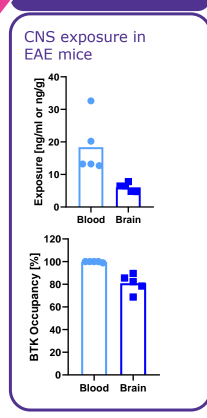
Evobrutinib targets inflammation and progression

Dual mechanism of action offers an innovative oral approach to MS therapy

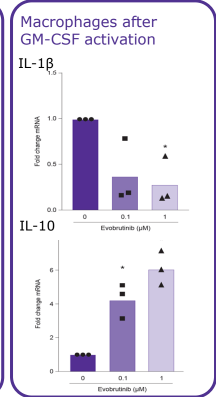
MOA on B-cells and macrophage/microglia cells is an innovative oral approach to MS therapy with potential to impact compartmentalized inflammation and progression



Evobrutinib is a CNS penetrant BTKi²

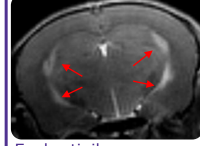


Evobrutinib shifts macrophage phenotypes from M1 pro- to M2 anti-inflammatory ³

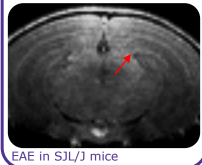


Evobrutinib reduces CNS B cells and leptomeningeal inflammation⁴





Evobrutinib:



- Important role of macrophage/ microglia cells in addition to B and T cells in both periphery and CNS¹
- Evobrutinib reaches brain to potentially impact compartmentalized inflammation²
- Effects may be mediated through effects on CNSresident and CNSmigrating cells^{3,4}



^{1.} Li et al. Nat Immunol 2018; 2. Adapted from Boschert U et al. ECTRIMS-ACTRIMS 2017; 3. Alankus YB et al. ECTRIMS 2018; 4. Sol Kim ECTRIMS 2020

Evobrutinib is optimally dosed to offer best-in-class BTK inhibition

Optimized dose selection & targeted covalent binding results in sustained BTK inhibition that is necessary for robust efficacy

Sustained BTK inhibition in CNS-resident and CNS-migrating cells resulting from covalent binding MOA and BID dosing can be critical to achieve best-in-disease efficacy¹

Targeted covalent binding leads to highly specific continuous target engagement²⁻⁶

Targeted covalent binding

Evobrutinib

Covalent binding

Tolebrutinib

Selectivity* Sustained action in CNS

Multiple other kinases

втк	Fenebrutinib	•

BID dosing enables critical >95% BTK inhibition throughout the day in the majority of patients⁷

BTK Occupancy (SS Trough)	25 mg QD	75 mg QD	75 mg BID
Threshold	% of P	opulation (RM	S Ph2)
95%	23	48	98
	No efficacy	Relapses	Maximum efficacy

Efficacy at 48 weeks ⁸			
75 mg QD Evobrutinib Fasted: ARR = 0.25	75 mg BID Evobrutinib Fasted: ARR = 0.11		

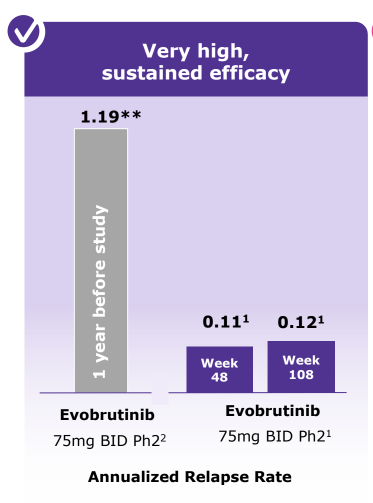


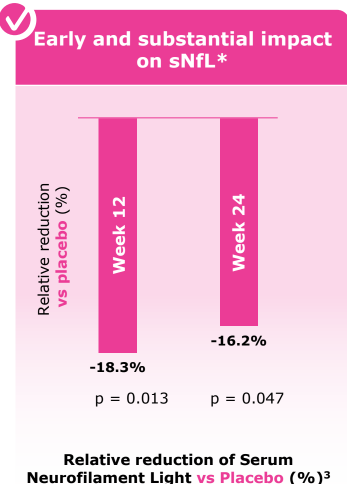
Reversible binding

Evobrutinib holds unmatched Long-Term Data among BTKi class in MS



Best-in-disease efficacy & favorable safety over 2 years in largest Phase II study in MS





- Evobrutinib is the only BTKi to have demonstrated very high, sustained efficacy and favorable safety in the largest Phase II study in MS (n=267), with an ARR of confirmed relapses of 0.12 up to 108 weeks¹
- Evobrutinib impacts sNfL levels, a biomarker of neuronal damage, reflecting disease activity and drug response in patients with MS³, starting at 12 weeks and maintained through 24 weeks²
- Evobrutinib is highly selective resulting in targeted kinase inhibition, and its safety data in >1200 patients over 2+ years supports the potential for an optimal longterm safety profile²

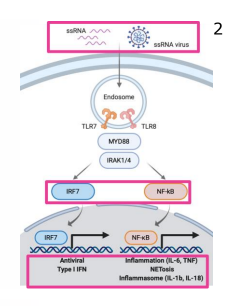


M5049 (TLR7/8 antagonist)

TLR7/8 are drivers of SLE pathology and possibly of COVID-19

Mechanism of Action¹

- M5049 (discovered in-house) is a potentially first-in class small molecule that blocks activation of Toll-like receptors TLR7 and TLR8, two innate immune sensors that detect single-stranded (ss) RNA from viruses such as SARS-COV-2, the virus responsible for COVID-19, and inflammatory self-RNAs in the context of autoimmunity
- Activation of TLR7/8 leads to immune cell activation and inflammation, which when not properly controlled can cause severe immunopathology



Results from
Phase I study in
healthy
volunteers
(NCT03676322)1

- Well-tolerated over the dosing interval, no significant or dose-limiting adverse event
- Pharmacokinetic parameters linear and dose-proportional from 1 to 200 mg
- Exposure-dependent inhibition of ex vivo-stimulated IL-6 secretion observed, with maximum inhibition achieved at 200 mg



Preliminary Phase I data warrant further investigation as a potential treatment for autoimmune diseases including SLE

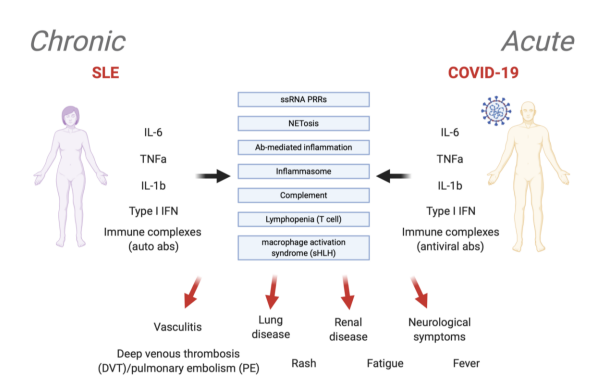
1: Port et al., A PHASE I, FIRST-IN-HUMAN STUDY TO ASSESS THE SAFETY, PHARMACOKINETICS AND PHARMACODYNAMICS OF SINGLE AND MULTIPLE ASCENDING DOSES OF M5049, A DUAL ANTAGONIST OF TLR7/8, IN HEALTHY SUBJECTS, Lupus Science & Medicine 2020;7(Suppl 1):A1-A131, conference cancelled due to COVID-19; 2 Adapted from ImmunoHorizons July 1, 2018 Dowling, D;

Acronyms: SLE = Systemic lupus erythematosus; TLR = Toll-like receptors

M5049 (TLR7/8 antagonist)

Similarities between SLE and COVID-19

Similarities between SLE and COVID-191



Phase II study started in July 2020

Rational:

- Investigate if M5049 intervention at critical point in course of COVID-19 disease may prevent or ameliorate hyperinflammatory response in patients with COVID-19 pneumonia and prevent progression to 'cytokine storm'
- Successful intervention with investigational drug may reduce life-threatening complications of COVID-19, including severe respiratory symptoms often necessitating further interventions such as mechanical ventilation

Design:

- Phase II randomized, controlled clinical study
- Commenced in July 2020

Results:



Dependent on recruitment and COVID-19 infection rates First results expected in Q3/Q4 2021

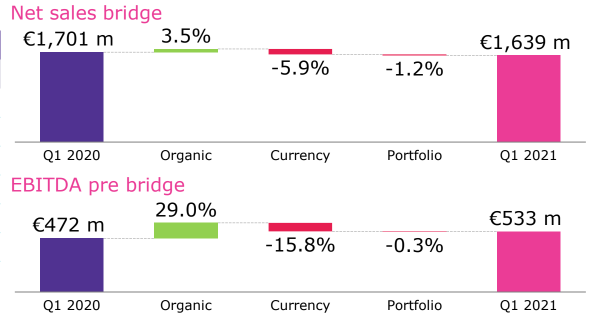


^{1:} Illustration created in-house; Acronyms: SLE = Systemic lupus erythematosus

Healthcare: strong Fertility & Oncology performance, while Mavenclad[®] remains impacted by depressed dynamic market

Healthcare P&L

[€m]	IFRS		Pre	
	Q1 2020	Q1 2021	Q1 2020	Q1 2021
Net sales	1,701	1,639	1,701	1,639
M&S [*]	-423	-370	-423	-365
Admin	-79	-73	-78	-69
R&D	-417	-416	-417	-415
EBIT	422	445	395	455
EBITDA	501	523	-	-
EBITDA pre	472	533	472	533
(in % of net sales)	27.8%	32.5%	27.8%	32.5%



Comments

- Mavenclad® growing +26 % organically to €147 m, amid still depressed dynamic market; Rebif® decline -17% in Q1 due to tough comps related to tender win in Q1 2020
- Oncology up +20%, Bavencio[®] sales doubled post UC 1L launch in U.S. and initial contribution from EU & JP¹; Erbitux[®] up +10% largely driven by China growth
- Base business about stable, strong Fertility growth (+22% org.) compensating for Rebif[®] and CM&E decline (-4% org.)

- Strong savings in M&S from continuous rigorous cost discipline, supported by reduced face-to-face activities amid pandemic
- R&D flat as a result of continued prioritization; no significant COVID-19 related project delays
- EBITDA pre and margin significantly supported by Bavencio[®] milestones (~ €50 m)



⁹⁰

¹ U.S., Europe and Japan approvals received on 30/06/2020, 25/01/2021 and 24/02/2021 respectively

Life science

Life Science

Capitalizing on three key life science trends



2 DIGITAL UNIVERSE

3 ASIA

Single Use / End to End

Opened Wuxi site in 2018, and expanded Danvers facility

Viral Vectors

Expanded Carlsbad viral vector manufacturing site in 2016; further doubling of capacity planned for 2021

Antibody Drug Conjugates (ADC)

Launched ADC Express[™] for the rapid production of ADCs #1 eCommerce site in Life Science¹

- > 90% of Millipore products on eCommerce platform
- x2 net sales growth of eCommerce vs.

Manufacturing/Distribution
Nantong, Wuxi Single use

Commercial expansion
Tier 2 cities

eCommerce partnership





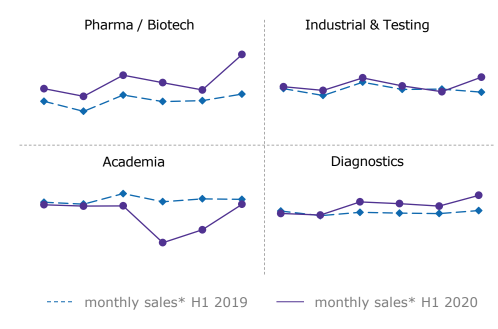
Life Science

Highly resilient and well positioned to participate in COVID-19 upside

2020 heatmap of COVID-19 impact by customer segment

Industrial & Pharma/ **Diagnostics** Academia Biotech Testing **Process** COVID-19 COVID-19 COVID-19 **Solutions** insensitive insensitive insensitive Research COVID-19 **Solutions** insensitive **Applied Solutions** downsides from COVID-19

H1 2020 monthly sales* by customer segment



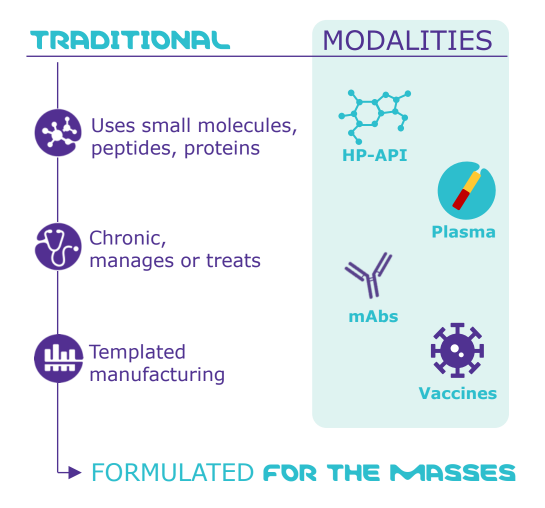
^{*} Ilustrative; to scale only within each customer segment

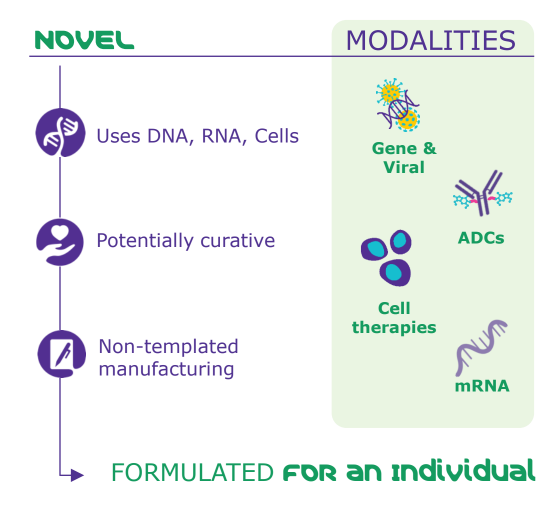


- Mid-term: downsides to fade, some upsides to stay, recovery started in June
- Order book for Process Solutions up by >40%, capacity expansion underway

Process Solutions: Therapies are evolving from treatments to cures

Advancing traditional is critical as novel modalities develop



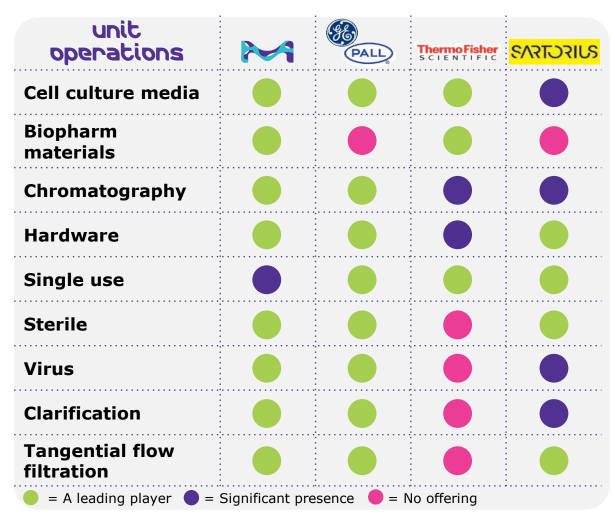


Acronyms: HP-API = highly potent active pharmaceutical ingredient; mAbs = monoclonal antibodies; DNA = deoxyribonucleic acid; (m)RNA = (messenger) ribonucleic acid; ADC = antibody drug conjugate



Process Solutions

COVID demands align with our strengths but increase supply chain pressure



COVID-19 Outlook

Type

Implications



65 programs

Bind and block virus from entering cells

- Universal templates
- A leading position for 8 out of 9 unit ops



Vaccine

199 programs
Protective **immune response**

- Multiple templates
- Leveraging Single Use



Nucleic Acid

43 programs
Leveraging human factory

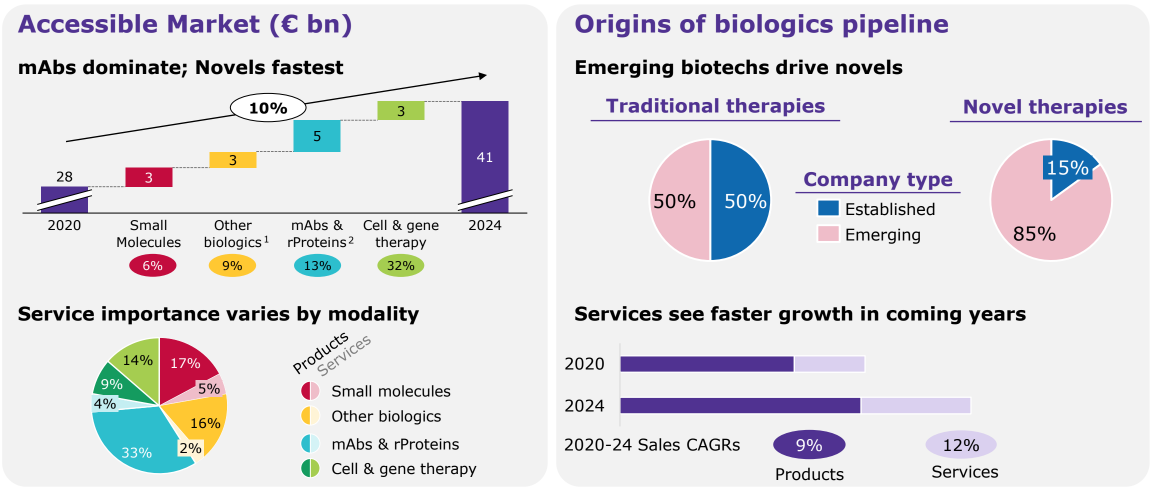
- Emerging manufacturing processes
- Lipids are critical

Sources: press releases, company reports, and internal assessments



Process Solutions

Opportunities in services to accelerate double-digit growth



Sources: Evaluate Pharma, internal market models, CSR sales data; ¹ Other biologics include plasma, vaccines, insulin, microbial and non-mAb biosimilars; ² mAbs include ADCs here; Additional acronym: rProteins = recombinant proteins



Process Solutions: Strategic direction

Innovate and invest today to continue above market growth in the future



Modality





ALL





NOVEL



Revenue driver



Near term



Mid term



Long term



Offering



Products

Filtration

Technology



Services



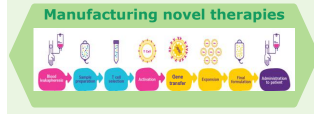
Products



Portfolio



- Contract development & manufacturing organization
- ✓ Contract testing organization
- ✓ Product characterization

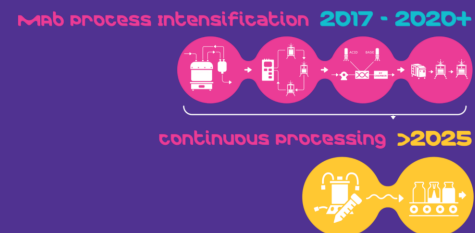


- Templated viral manufacturing consumables and reagents
- ✓ Autologous manufacturing system



Next-generation bioprocessing on the cards



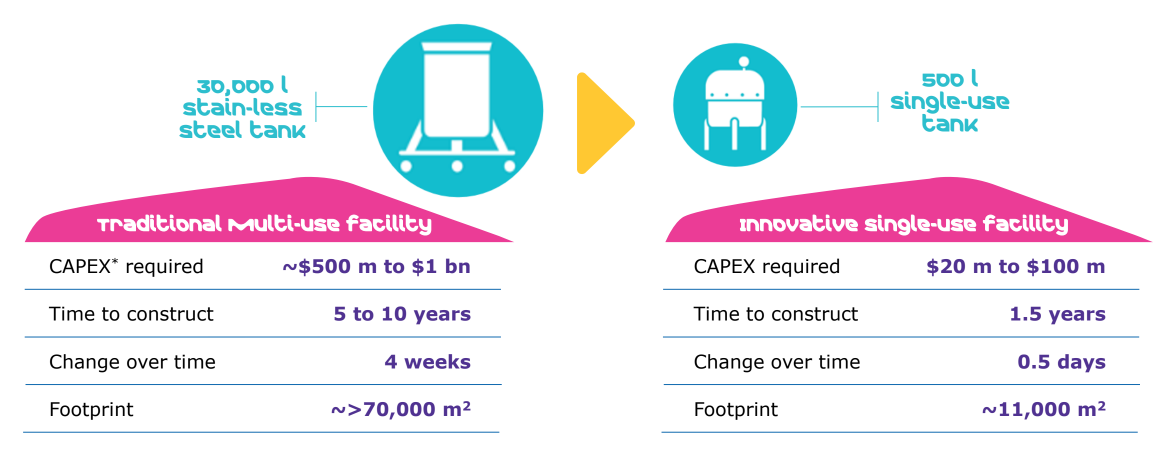


Continuous bioprocessing will ...

- be an evolution in mAb bioprocessing
- take time to establish
- leverage the present
- lead to hybrid solutions

Process Solutions

Our single-use technologies drive flexibility in modern bioprocessing





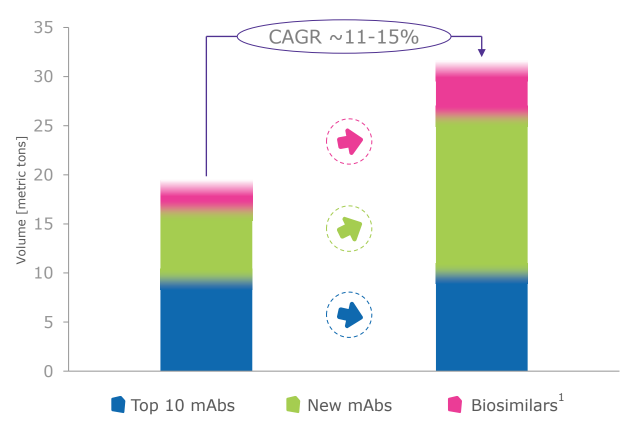
Strong demand for single-use technologies and Process Solutions' broad offering was and will remain a key source of growth for Life Science



Process Solutions

mAbs market democratization will drive diversification, change & variability

mAb volume projections 2020 to 2024



¹Biosimilars scaling factor = 2.8 based off internal estimates and McKinsey analysis; Source: company estimate based on industry forecasts, EvaluatePharma; mAbs = Monoclonal antibodies

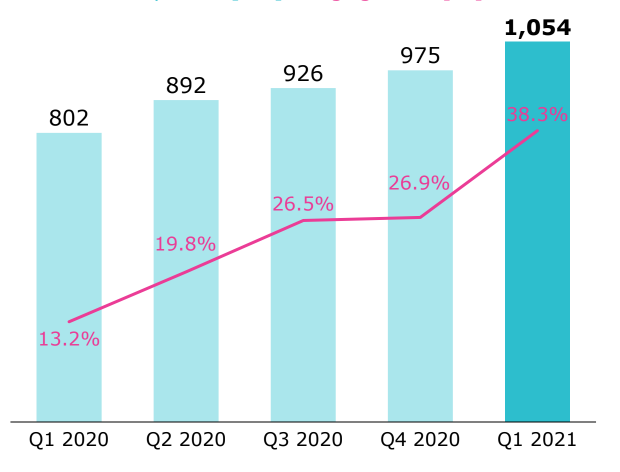
market development

- Overall mAbs market will grow ~11-15% CAGR
- Top 10 originator mAbs represent
 50% of market volume today and will decline to ~30% in 2024
- Biosimilars will gain share



BIG 3 - Process Solutions: upside potential continues to materialize

Sales development [€m] - org. growth [%]

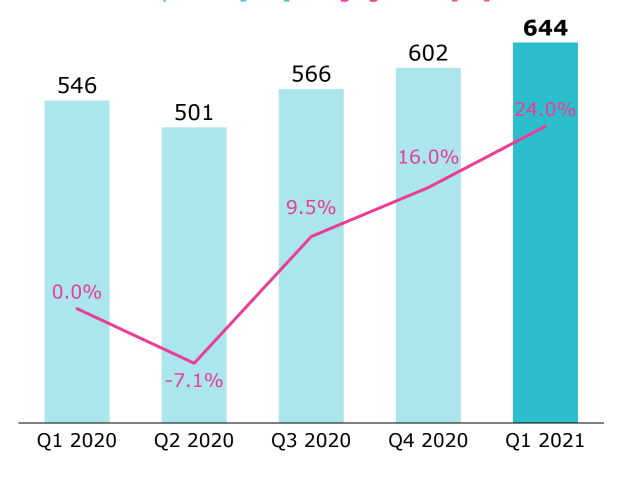


- COVID-19 related projects in BioP Single Use
 and Downstream remain key drivers of
 50% additional pandemic-related growth
- Unprecedented growth across all regions particularly strong in **Asia** (>50% org.) in large parts driven by the recovery vs. Q1 2020 lockdown in China
- Pharma & Biotech customer segment remains by far the strongest growth driver
- **Comps will be getting tough** from Q2 2021 onwards, as base starts including significant COVID-19 business upside
- FX headwinds of -7% mute absolute numbers



Research Solutions: additional COVID-19 demand has gained momentum

Sales development [€m] - org. growth [%]



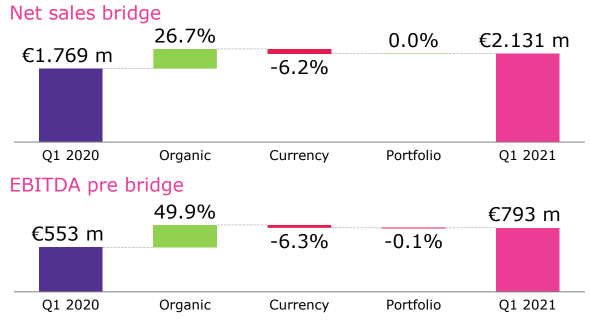
- Ongoing core business recovery boosted by additional COVID-19 related demand
- Unprecedented growth across all regions particularly strong in **Asia** (>30% org.) and **North America** (>20% org.)
- Pharma & Biotech customer segment strongest Q1 growth driver in absolute terms;
 Diagnostics customer segment showed fastest growth in relative terms
- Q2 2021 will be softest comp, against the COVID-19 induced lock-down dip in 2020
- FX headwinds of -6% mute absolute numbers



Life Science: strong core business and rising COVID-19 demand fuel record growth, particularly in Process and Research Solutions

Life Science P&L

[€m]	IFRS		Pr	е
	Q1 2020	Q1 2021	Q1 2020	Q1 2021
Net sales	1,769	2,131	1,769	2,131
M&S*	-498	-501	-497	-500
Admin	-89	-82	-80	-74
R&D	-75	-75	-75	-75
EBIT	345	593	357	607
EBITDA	541	779	_	-
EBITDA pre	553	793	553	793
(in % of net sales)	31.2%	37.2%	31.2%	37.2%



Comments

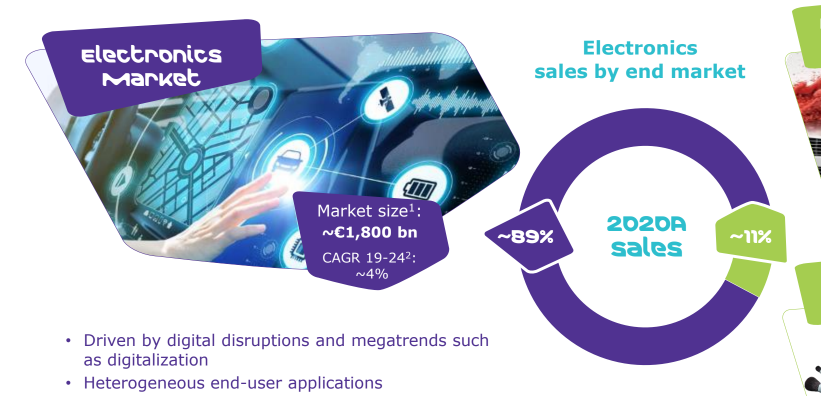
- Process Solutions org. growth of +38% driven by downstream & single use for COVID-19 projects; strong underlying demand supported by acceleration of capacity expansions
- Research Solutions speeding up further to an exceptional +24% organic growth, driven by recovery in base business and additional COVID-19 opportunities, mainly in diagnostics and pharma
- Applied Solutions growth (+8% org.) driven by APAC recovery against low Q1 2020

- Declining M&S in % of sales from 28% to 23% despite higher logistics cost amid pandemic
- Flat R&D spend with continued focused investments in strategic projects in high growth & emerging segments
- Business performance, operational leverage & favorable mix continue to drive strong EBITDA pre and margin expansion despite 6% FX drag



Electronics

Electronics targets attractive markets – especially in the electronics space



Automotive Market



 Increasing demand in emerging markets

Market size: ~€2,000 bn

cosmetics market

- Driven by world GDP growth
- Rising living standards and higher disposable income

Market size: ~€400 bn

• Semiconductors as the engine of all electronic

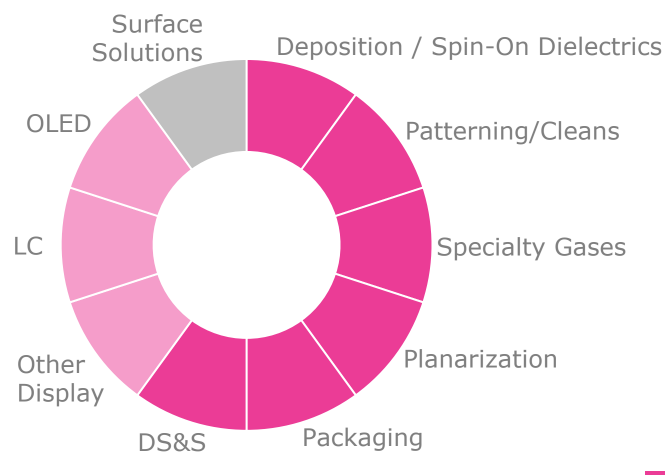


systems

¹Prismark 2021; ² Internal estimation

Electronics

Expected mid-term portfolio split



Mid-term the Electronics portfolio will consist of ~10 equally sized businesses

>60% of them serving chip makers

Juliace Joidholls

 ${\tt INDICATIVE\ Chart}$

Semiconductor Solutions

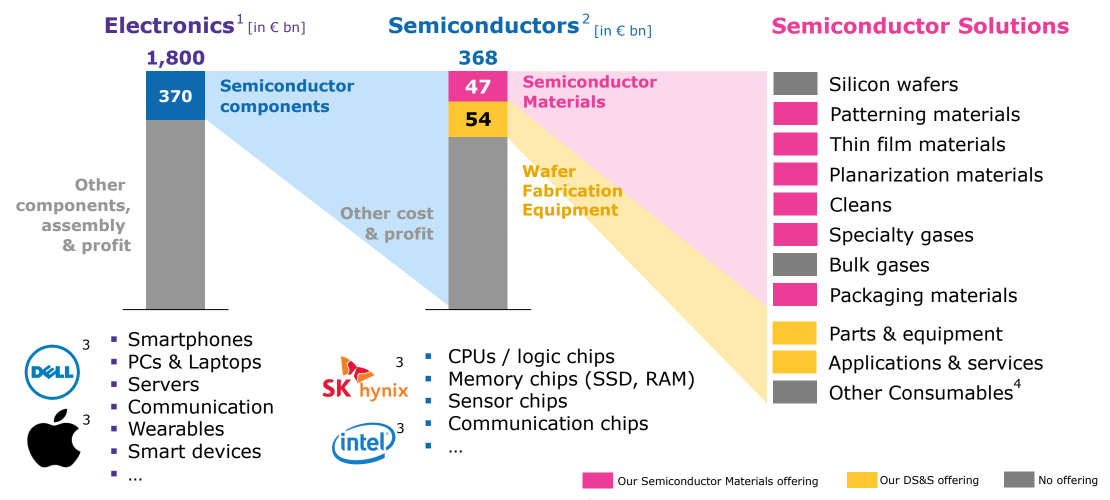


Display Solutions



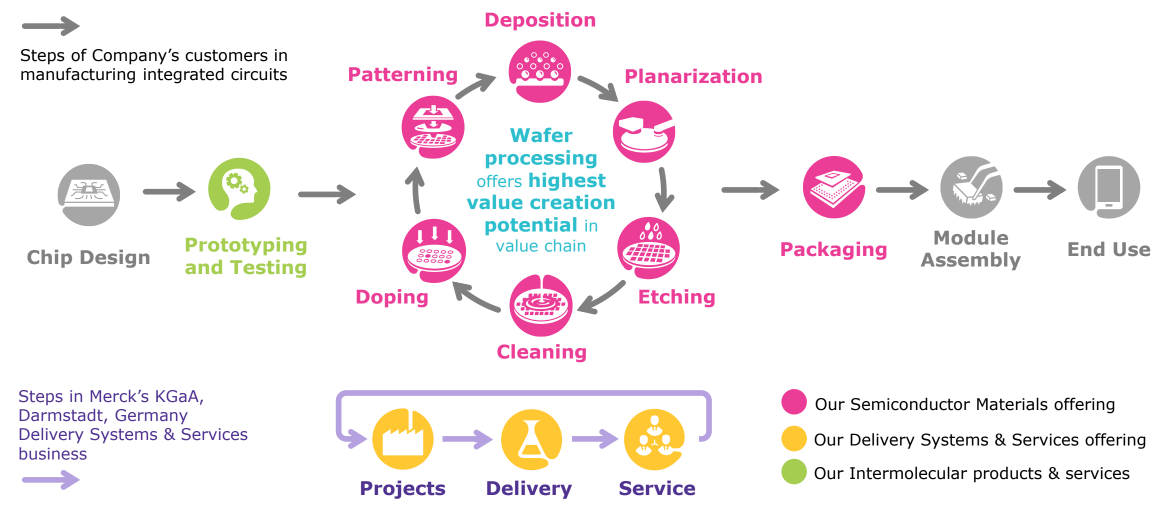


Semiconductor Solutions - integrated materials player, well positioned to serve the need of customers in semiconductor fabrication



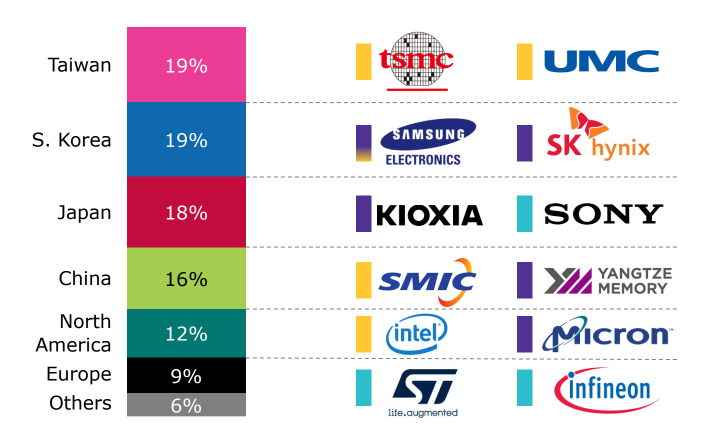
llustrative Industry P&Ls based on Sources: ¹Prismark 2021, ²Prismark 2021 & WSTS/SIA & SEMI Q1 2020; ³Representative player in the industry, non-exclusive list, not based on any underlying criteria; ⁴e.g. Filters, Pads, etc.; CPU = Central Processing Unit; RAM = Random Access Memory; SSD = Solid State Disk; CMOS = Complementary metal-oxide semiconductor

Unique comprehensive products and services portfolio offers end-to-end solutions, well-placed in high growth segments



Beyond a comprehensive portfolio Semiconductor Solutions also has an industry spanning customer base, supplying various end markets

2019 wafer capacity by region Selected customers per region 2



Semiconductor Solutions has

customers

supplying all top 10 chip makers and virtually all of the top 100³



Focus on logic chips

Focus on memory chips

Focus on other chips

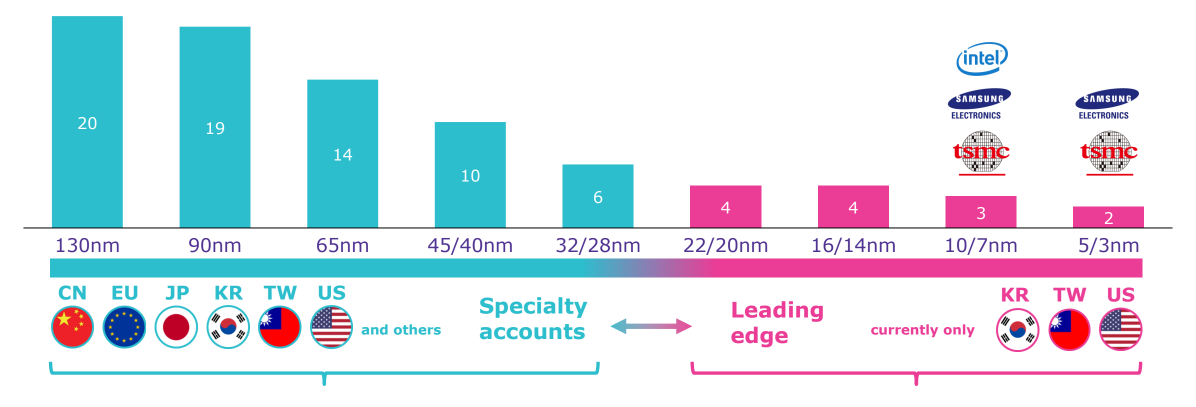
¹SEMI World Fab Forecast Q3 2020 - Dec 2019 capacity,

Representative, non-exclusive list, not based on any underlying criteria

³Based on H1 2020 Sales

Only 3 companies are currently running volume production ≤10nm These companies have the largest market shares across all nodes

Number of companies currently running volume production per logic chip node¹



Primary focus on operation efficiency

Primary focus on innovation efficiency

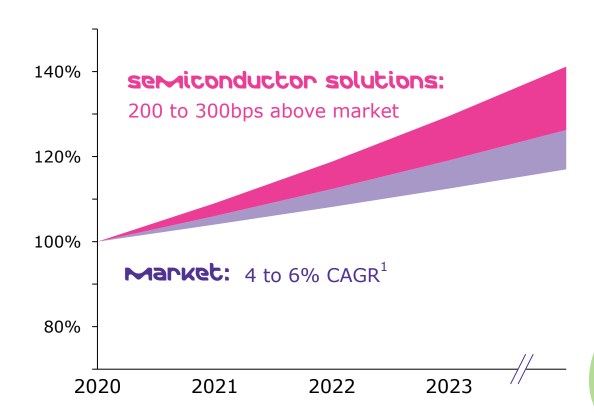
¹Source: Wikichip.org and own data; volume production as of Sep 2020; countries are listed in alphabetical order

Semiconductor Materials

Set to outgrow highly attractive semiconductor materials market

Semiconductor Solutions sales guidance vs. market

[Indexed 2020 = 100%]



¹Source: Jan 2020 IC Insights 2018-2024 CAGR for wafer starts in million units

Market

- Technological trends inevitably drive exponential data growth
- More data requires more chips and higher complexity of chips
- Rising materials value added per wafer

semiconductor solutions

- Comprehensive offering focusing on attractive materials categories
- Integration topline synergies
- Critical mass and deep customer centricity
- Better customer understanding: know-how exchange and collaboration across DS&S and Semiconductor Materials
- Cutting-edge innovation and R&D capabilities



Technology trends inevitably drive exponential data growth... ...more chips needed to generate, transfer, process & store data

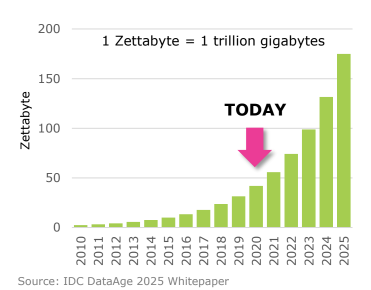
Data created worldwide

is growing +30% annually

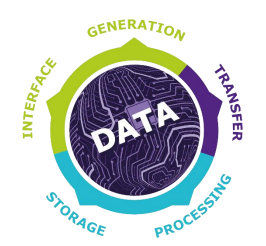
All segments of data application are affected by global data growth

Technology trends strongly impact relevance of data application segments

Size of global data sphere









Technology market growth - examples

5G Technology¹

>122% CAGR

Artificial Intelligence²

>33% CAGR

IoT Sensors³

>24% CAGR

Data Center Services⁴

>13% CAGR

Autonomous Driving⁵

>18% CAGR

Semiconductor Solutions supports growth trend as part of "the company behind the companies, advancing digital living"

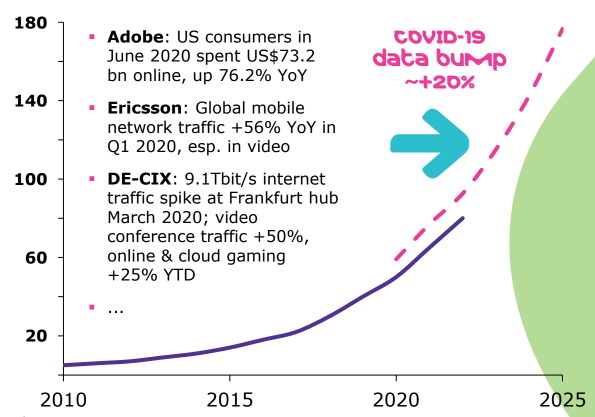
1) alliedmarketresearch.com, Prismark 2020, CAGR 2021-2026; 2) fortunebusinessinsights.com, post-gazette.com, CAGR 2018-2026; 3) mordorintelligence.com, computerweekly.com, CAGR 2020-2025; 4) mordorintelligence.com, Prismark 2020; CAGR 2020-2025; 5G = 5th-generation cellular wireless; IoT = Internet of Things 5) mordorintelligence.com, autonomous car market value CAGR 2020-2025



Semiconductor Solutions

COVID-19 has vaulted the "digital transformation" by ~5 years¹

Expected COVID-19 impact on global datasphere² [zetabytes]



¹Source: McKinsey May 2020 "The COVID-19 recovery will be digital: A plan for the first 90 days"; ²Source: Seagate, IDC April 2020, Merck KGaA, Darmstadt, Germany

covid-19 impact on data growth expected to be positive

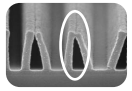
- Work-from-home/stay-at-home economy
 - Significant increase in video conferences, online shopping, online gaming, streaming
- Change in consumers' and enterprises' digital behavior expected to be long-lasting
- Need for more, faster & more reliable data processing, storage and bandwidth
 - Acceleration of semiconductor demand

semiconductor solutions stands ready to support increased demand



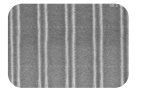
Expanding the limits of how small you can go

Pattern collapse



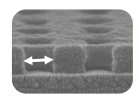


Lithography limitation



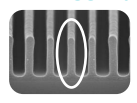


Wide features





AZ® rinse materials



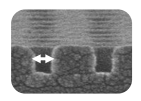
As lines get narrower and closer together in advanced chip generation, lines tend to "stick" due to surface tension.

Directed self-assembly (DSA)



Block copolymer can generate small lines or contact holes by selfassembly. This allows miniaturization without expensive new equipment.

AZ® shrink materials



Shrink materials "shrink" the gap between lines and, hence, allow the manufacture of narrower features otherwise not possible.



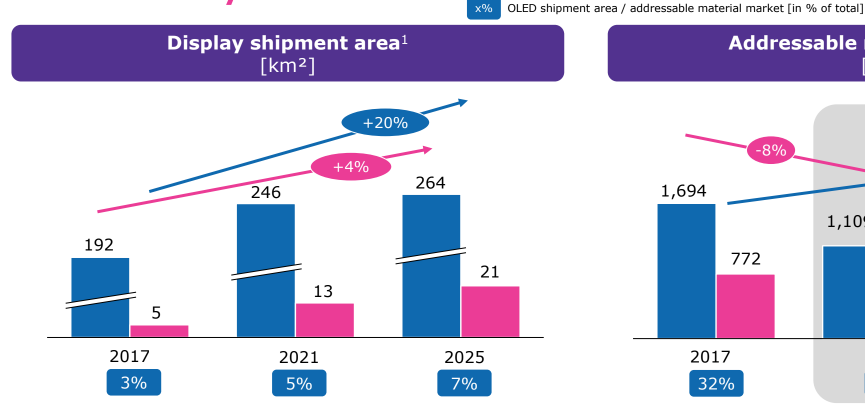
Group delivers highly innovative solutions for complex customer problems



Electronics

Display Solutions - OLED material market to exceed LC material

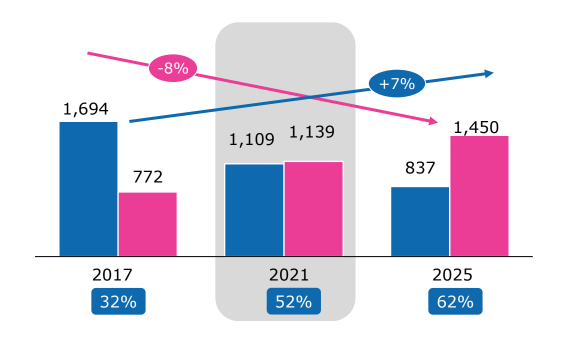
market already in 2021



- Continued growth across all technologies
- OLED growing faster than LCD, but LCD to command 90+% area share for foreseeable future

Addressable material market² [€m]

Liquid Crystals



- Material value per OLED display higher than in LCD
- OLED material market to exceed LC material market by 2021, but market split between many more players



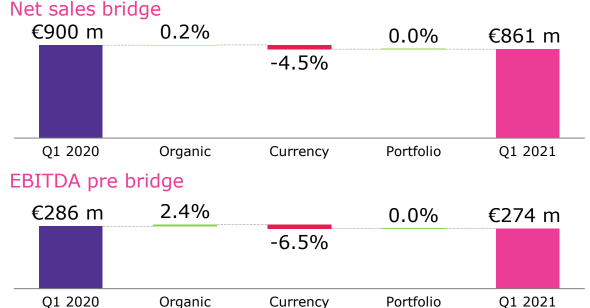


OLED

Electronics: year-over-year and vs. Q4 2020 Semi growth as well as recovering Surface compensate Display Solutions decline

Electronics P&L

[€m]	IFRS		Pr	e
	Q1 2020	Q1 2021	Q1 2020	Q1 2021
Net sales	900	861	900	861
M&S [*]	-136	-135	-134	-135
Admin	-38	-34	-38	-33
R&D	-71	-67	-73	-66
EBIT	116	126	151	142
EBITDA	251	260	-	-
EBITDA pre	286	274	286	274
(in % of net sales)	31.7%	31.8%	31.7%	31.8%



Comments

- Semiconductor Solutions: solid Semiconductor Materials growth across all product categories muted by DS&S project launch phasing; continued confidence in strong outlook for FY 2021
- Display Solutions: down -7% organically as OLED growth not yet compensating for Liquid Crystals decline
- Surface Solutions: returning to 5% organic growth, mainly supported by recovery in the automotive industry

- Stable M&S despite higher logistic costs, while Admin and R&D are declining
- All P&L lines continue to reflect diligent cost management amid Bright Future transformation and Versum integration synergies
- EBITDA pre (+2% org.) exceeds sales growth but burdened by -7% FX headwinds



CONSTANTIN FEST



Head of Investor Relations +49 6151 72-5271 constantin.fest@emdgroup.com

SVENJA BUNDSCHUH



Assistant Investor Relations +49 6151 72-3744 svenja.bundschuh@emdgroup.com

SARA HOFMANN



Assistant Investor Relations +49 6151 72-3321 sara.hofmann@emdgroup.com

ILJA DOERING



Institutional Investors / Analysts +49 6151 72-24164 ilja.doering@emdgroup.com

GUNNAR ROMER



Institutional Investors /
Analysts
+49 6151 72-2584
gunnar.romer@emdgroup.com

AMELIE SCHRADER



Institutional Investors /
Analysts
+49 6151 72-22076
amelie.schrader@emdgroup.com

EVA STERZEL



ESG / Institutional & Retail Investors / AGM +49 6151 72-5355 eva.sterzel@emdgroup.com



EMAIL: <u>investor.relations@emdgroup.com</u>

WEB: <u>www.emdgroup.com/investors</u>

FAX: +49 6151 72-913321