

GLAXOSMITHKLINE PLC

Form 20-F

March 16, 2018

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As filed with the Securities and Exchange Commission on March 16, 2018

UNITED STATES

SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 20-F

**REGISTRATION STATEMENT PURSUANT TO SECTION 12(b) OR (g) OF THE SECURITIES
EXCHANGE ACT OF 1934**

OR

**ANNUAL REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT
OF 1934**

For the fiscal year ended December 31, 2017

OR

**TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE
ACT OF 1934**

OR

**SHELL COMPANY REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES
EXCHANGE ACT OF 1934**

Commission file number 1-15170

GlaxoSmithKline plc

(Exact name of Registrant as specified in its charter)

England

(Jurisdiction of incorporation or organization)

980 Great West Road, Brentford, Middlesex TW8 9GS England

(Address of principal executive offices)

Victoria Whyte

Company Secretary

GlaxoSmithKline plc

980 Great West Road

Brentford, TW8 9GS

England

+44 20 8047 5000

company.secretary@gsk.com

(Name, Telephone, E-mail and/or Facsimile number and Address of Company Contact Person)

Securities registered or to be registered pursuant to Section 12(b) of the Act:

Title of Each Class	Name of Each Exchange On Which Registered
American Depositary Shares, each representing	
2 Ordinary Shares, Par value 25 pence	New York Stock Exchange
5.650% Notes due 2018	New York Stock Exchange
2.850% Notes due 2022	New York Stock Exchange
2.800% Notes due 2023	New York Stock Exchange

5.375% Notes due 2034

London Stock Exchange

6.375% Notes due 2038

New York Stock Exchange

4.200% Notes due 2043

New York Stock Exchange

Securities registered or to be registered pursuant to Section 12(g) of the Act:

None

(Title of class)

Securities for which there is a reporting obligation pursuant to Section 15(d) of the Act:

None

(Title of class)

Indicate the number of outstanding shares of each of the issuer's classes of capital or common stock as of the close of the period covered by the annual report.

Ordinary Shares of Par value 25 pence each

5,372,553,820

Indicate by check mark if the registrant is a well-known seasoned issuer, as defined in Rule 405 of the Securities Act.

Yes No

If this report is an annual or transition report, indicate by check mark if the registrant is not required to file reports pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934.

Yes No

Note: Checking the box above will not relieve any registrant required to file reports pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934 from their obligations under those Sections.

Indicate by check mark whether the registrant (1) has filed all reports to be filed by Section 13 or 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period that the registrant was required to file such reports), and (2) has been subject to such filing requirements for the past 90 days.

Yes No

Indicate by check mark whether the registrant has submitted electronically and posted on its corporate Web site, if any, every Interactive Data File required to be submitted and posted pursuant to Rule 405 of Regulation S-T (§232.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit and post such files).

Yes No

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Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, or a non-accelerated filer. See definition of "accelerated filer" and "large accelerated filer" in Rule 12b-2 of the Exchange Act. (Check one):

Large accelerated filer

Accelerated filer
company

Non-accelerated filer

Emerging growth

If an emerging growth company that prepares its financial statements in accordance with U.S. GAAP, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

The term "new or revised financial accounting standard" refers to any update issued by the Financial Accounting Standards Board to its Accounting Standards Codification after April 5, 2012.

Indicate by check mark which basis of accounting the registrant has used to prepare the financial statements included in this filing:

U.S. GAAP

International Financial Reporting Standards as issued

Other

by the International Accounting Standards Board

If "Other" has been checked in response to the previous question, indicate by check mark which financial statement item the registrant has elected to follow.

Item 17 Item 18

If this is an annual report, indicate by check mark whether the registrant is a shell company (as defined in Rule 12b-2 of the Exchange Act).

Yes No

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* As permitted by Rule 405(a)(2)(ii) of Regulation S-T, the registrant's XBRL (eXtensible Business Reporting Language) information will be furnished in an amendment to this Form 20-F that will be filed no more than 30 days after the date hereof. In accordance with Rule 402 of Regulation S-T, the information in these exhibits shall not be deemed to be filed for purposes of Section 18 of the Exchange Act, or otherwise subject to the liability of that section, and shall not be incorporated by reference into any registration statement or other document filed under the Securities Act, or the Exchange Act, except as shall be expressly set forth by specific reference in such filing.

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Pursuant to Rule 12b-23(a) of the Securities Exchange Act of 1934, as amended, the information for GlaxoSmithKline plc's Form 20-F for the year ended December 31, 2017 as set out below is being incorporated by reference from the GSK Annual Report 2017 included as exhibit 15.3 to this Form 20-F dated and submitted on March 16, 2018 (the GSK Annual Report 2017).

All references in this Form 20-F to GlaxoSmithKline, the Group, GSK, we or our mean GlaxoSmithKline plc and its subsidiaries; the company means GlaxoSmithKline plc.

References below to major headings include all information under such major headings, including subheadings, unless such reference is a reference to a subheading, in which case such reference includes only the information contained under such subheading.

In addition to the information set out below, the information set forth under the headings Cautionary statement on the inside back cover, Directors Report on page 112, Directors statement of responsibilities on page 148, Directors statement of responsibilities in relation to the company's financial statements on page 233, Share capital and control on pages 267 to 268, Financial calendar, Results announcements and Financial reports on page 269, Annual General Meeting 2018 on page 270, Registrar on page 272 ADS Depositary, Glaxo Wellcome and SmithKline Beecham Corporate PEPs, Donating shares to Save the Children, Contacts, Share scam alert and Responsible Business Supplement on page 273 Section 13(r) of the US Securities Exchange Act on page 275 and Glossary of terms on page 287 in each case of the GSK Annual Report 2017 is incorporated by reference.

Notice regarding limitations on Director Liability under English Law

Under the UK Companies Act 2006, a safe harbour limits the liability of Directors in respect of statements in and omissions from certain portions of the GSK Annual Report 2017 incorporated by reference herein, namely the Directors Report (for which see page 112 thereof), the Strategic Report (pages 1 to 78 thereof, portions of which are incorporated by reference as described below) and the Remuneration Report (pages 113 to 146 portions of which are incorporated by reference as described below). These reports have been drawn up and presented in accordance with, and in reliance upon, English company law. Under English law, the Directors would be liable to the company, but not to any third party, if these sections of the GSK Annual Report 2017 contain errors as a result of recklessness or knowing misstatement or dishonest concealment of a material fact, but would not otherwise be liable.

Portions of the GSK Annual Report 2017 incorporated by reference herein contain references to our website. Information on our website or any other website referenced in the GSK Annual Report 2017 is not incorporated into this Form 20-F and should not be considered to be part of this Form 20-F. We have included any website as an inactive textual reference only.

PART I

Item 1. **Identity of Directors, Senior Management and Advisers**
Not applicable.

Item 2. **Offer Statistics and Expected Timetable**
Not applicable.

Item 3. **Key Information**

3.A Selected financial data

The information set forth under the heading:

Five year record on pages 248 to 250; and

Dividends on page 269
of the GSK Annual Report 2017 is incorporated herein by reference.

3.B Capitalization and indebtedness

Not applicable.

3.C Reasons for the offer and use of proceeds

Not applicable.

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3.D Risk Factors

Principal risks and uncertainties

The principal risks discussed below are the risks and uncertainties relevant to our business, financial condition and results of operations that may affect our performance and ability to achieve our objectives. The risks below are those that we believe could cause our actual results to differ materially from expected and historical results.

We must adapt to and comply with a broad range of laws and regulations. These requirements apply to research and development, manufacturing, testing, approval, distribution, sales and marketing of Pharmaceutical, Vaccine and Consumer Healthcare products and affect not only the cost of product development but also the time required to reach the market and the likelihood of doing so successfully.

Moreover, as rules and regulations change, and governmental interpretation of those rules and regulations evolves, the nature of a particular risk may change. Changes to certain regulatory regimes may be substantial. Any change in, and any failure to comply with, applicable law and regulations could materially and adversely affect our financial results.

Similarly, our global business exposes us to litigation and government investigations, including but not limited to product liability litigation, patent and antitrust litigation and sales and marketing litigation. Litigation and government investigations, including related provisions we may make for unfavourable outcomes and increases in related costs such as insurance premiums, could materially and adversely affect our financial results.

More detail on the status and various uncertainties involved in our significant unresolved disputes and potential litigation is set out in Note 45, Legal proceedings, on pages 227 to 232 of the GSK Annual Report 2017.

UK regulations require a discussion of the mitigating activities a company takes to address principal risks and uncertainties. A summary of the activities that the Group takes to manage each of our principal risks accompanies the description of each principal risk below. The principal risks and uncertainties are not listed in order of significance.

Patient safety

Risk definition

Failure to appropriately collect, review, follow up, or report adverse events from all potential sources, and to act on any relevant findings in a timely manner.

Risk impact

The risk impact has the potential to compromise our ability to conduct robust safety signal detection and interpretation and to ensure that appropriate decisions are taken with respect to the risk/benefit profile of our products, including the completeness and accuracy of product labels and the pursuit of additional studies/analyses, as appropriate. This could lead to potential harm to patients, reputational damage, product liability claims or other litigation, governmental investigation, regulatory action such as fines, penalties or loss of product authorisation.

Context

Pre-clinical and clinical trials are conducted during the development of investigational Pharmaceutical, Vaccine and Consumer Healthcare products to determine the safety and efficacy of the products for use by humans. Notwithstanding the efforts we make to determine the safety of our products through appropriate pre-clinical and

clinical trials, unanticipated side effects may become evident only when products are widely introduced into the marketplace. Questions about the safety of our products may be raised not only by our ongoing safety surveillance and post-marketing studies but also by governmental agencies and third parties that may analyse publicly available clinical trial results.

The Group is currently a defendant in a number of product liability lawsuits, including class actions, that involve significant claims for damages related to our products. Litigation, particularly in the US, is inherently unpredictable. Class actions that seek to sweep together all persons who take our products increase the potential liability. Claims for pain and suffering and punitive damages are frequently asserted in product liability actions and, if allowed, can represent potentially open-ended exposure and thus, could materially and adversely affect the Group's financial results.

Product quality

Risk definition

Failure to comply with current Good Manufacturing Practices (cGMP) or inadequate controls and governance of quality in the supply chain covering supplier standards, manufacturing and distribution of products.

Risk impact

A failure to ensure product quality could have far reaching implications in terms of patient and consumer safety resulting in product launch delays, supply interruptions and product recalls. This would have the potential to do damage to our reputation, as well as result in other regulatory, legal and financial consequences.

Context

Patients, consumers and HCPs trust the quality of our products. Product quality may be influenced by many factors including product and process understanding, consistency of manufacturing components, compliance with GMP, accuracy of labelling, reliability of the external supply chain, and the embodiment of an overarching quality culture. The internal and external environment continues to evolve as new products and new legislation are introduced. Critically, we are addressing the impact of Brexit on our supply chain management and quality oversight between the UK and the EU and are developing and deploying appropriate contingency plans to avoid interruption of supply to patients.

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Financial controls and reporting

Risk definition

Failure to comply with current tax laws or incurring significant losses due to treasury activities; failure to report accurate financial information in compliance with accounting standards and applicable legislation.

Risk impact

Non-compliance with existing or new financial reporting and disclosure requirements, or changes to the recognition of income and expenses, could expose us to litigation and regulatory action and could materially and adversely affect our financial results. Changes in tax laws or in their application with respect to matters such as transfer pricing, foreign dividends, controlled companies, R&D tax credits, taxation of intellectual property or a restriction in tax relief allowed on the interest on debt funding, could impact our effective tax rate. Significant losses may arise from inconsistent application of treasury policies, transactional or settlement errors, or counterparty defaults.

Any changes in the substance or application of the governing tax laws, failure to comply with such tax laws or significant losses due to treasury activities could materially and adversely affect our financial results.

Context

The Group is required by the laws of various jurisdictions to disclose publicly its financial results and events that could materially affect the financial results of the Group. Regulators routinely review the financial statements of listed companies for compliance with new, revised or existing accounting and regulatory requirements. The Group believes that it complies with the appropriate regulatory requirements concerning our financial statements and disclosure of material information including any transactions relating to business restructuring such as acquisitions and divestitures. However, should we be subject to an investigation into potential non-compliance with accounting and disclosure requirements, this may lead to restatements of previously reported results and significant penalties.

Our Treasury group deals in high value transactions, mostly foreign exchange and cash management transactions, on a daily basis. These transactions involve market volatility and counterparty risk. The Group's effective tax rate reflects rates of tax in the jurisdictions in which the Group operates that are both higher and lower than the UK rate and takes into account regimes that encourage innovation and investment in science by providing tax incentives which, if changed, could affect the Group's tax rate. In addition, the worldwide nature of our operations means that our intellectual property, R&D and manufacturing operations are centred in a number of key locations. A consequence of this is that our cross-border supply routes, necessary to ensure supplies of medicines into numerous end markets, can be complex and result in conflicting claims from tax authorities as to the profits to be taxed in individual countries. Tax legislation itself is also complex and differs across the countries in which we operate. As such, tax risk can also arise due to differences in the interpretation of such legislation. The tax charge included in our financial statements is our best estimate of tax liability pending audits by tax authorities.

We expect there to be continued focus on tax reform in 2018 and future years driven by the Organisation for Economic Cooperation & Development's Base Erosion and Profit Shifting project and European Commission initiatives including the use of fiscal state aid investigations. Together with domestic initiatives around the world, these may result in significant changes to established tax principles and an increase in tax authority disputes. These, regardless of their merit or outcomes, can be costly, divert management attention and may adversely impact our reputation and relationship with key stakeholders.

Anti-bribery and corruption

Risk definition

Failure of GSK employees, consultants and third parties to comply with our Anti-bribery & corruption (ABAC) principles and standards, as well as with all applicable legislation.

Risk impact

Failure to mitigate this risk could expose the Group and associated persons to governmental investigation, regulatory action and civil and criminal liability and may compromise the Group's ability to supply its products under certain government contracts. In addition to legal penalties, a failure to prevent bribery through complying with ABAC legislation and regulations could have substantial implications for the reputation of the company, the credibility of senior leaders, and an erosion of investor confidence in our governance and risk management.

Context

We are exposed to bribery and corruption risk through our global business operations. In some markets, the government structure and the rule of law are less developed, and this has a bearing on our bribery and corruption risk exposure. In addition to the global nature of our business, the healthcare sector by its very nature maintains relationships with government bodies, is highly competitive and subject to regulation. This increases the instances where we are exposed to activities and interactions with bribery and corruption risk.

The Group has been subject to a number of ABAC inquiries. We reached a resolution with the US authorities in 2016 regarding their ABAC inquiry, following which we are subject to a self-monitoring arrangement until September 2018. Government investigations regarding our China and other business operations are ongoing. These investigations are discussed further in Note 45, Legal proceedings.

Commercial practices

Risk definition

Failure to engage in commercial activities that are consistent with the letter and spirit of legal, industry, or the Group's requirements relating to marketing and communications about our medicines and associated therapeutic areas; appropriate interactions with HCPs and patients; and legitimate and transparent transfer of value.

Risk impact

Failure to manage risks related to commercial practices could materially and adversely affect our ability to grow a diversified global business and deliver more products of value for patients and consumers. Failure to comply with applicable laws, rules and regulations may result in governmental investigation, regulatory action and legal proceedings brought against the Group by governmental and private plaintiffs which could result in government sanctions, and criminal and/or financial penalties. Failure to provide accurate and complete information related to our products may result in incomplete awareness of the risk/benefit profile of our products and possibly suboptimal treatment of patients and consumers.

Any practices that are found to be misaligned with our values could also result in reputational harm and dilute trust established with external stakeholders.

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Context

We operate on a global basis in an industry that is both highly competitive and highly regulated. Our competitors may make significant product innovations and technical advances and may intensify price competition. In light of this competitive environment, continued development of commercially viable new products and the development of additional uses for existing products that reflect insights which help ensure those products address the needs of patients/consumers, HCPs, and payers are critical to achieve our strategic objectives.

As do other pharmaceutical, vaccine and consumer companies, we face downward price pressure in major markets, declining emerging market growth, and negative foreign exchange impact.

Developing new Pharmaceutical, Vaccine and Consumer Healthcare products is a costly, lengthy and an uncertain process. A product candidate may fail at any stage, including after significant economic and human resources have been invested. Our competitors' products or pricing strategies or any failure on our part to develop commercially successful products, or to develop additional uses for existing products, could materially and adversely affect our ability to achieve our strategic objectives.

We are committed to the ethical and responsible commercialisation of our products to support our mission to improve the quality of human life by enabling people to do more, feel better, and live longer. To accomplish this mission, we engage the healthcare community in various ways to provide important information about our medicines. Promotion of approved products seeks to ensure that HCPs globally have access to information they need, that patients and consumers have access to the information and products they need and that products are prescribed, recommended or used in a manner that provides the maximum healthcare benefit to patients and consumers. We are committed to communicating information related to our approved products in a responsible, legal, and ethical manner.

Research practices

Risk definition

Failure to adequately conduct ethical and sound preclinical and clinical research. In addition, failure to engage in scientific activities that are consistent with the letter and spirit of the law, industry, or the Group's requirements, and failure to secure adequate patent protection for GSK's products.

Risk impact

The impacts of the risk include harm to human subjects, reputational damage, failure to obtain the necessary regulatory approvals for our products, governmental investigation, legal proceedings brought against the Group by governmental and private plaintiffs (product liability suits and claims for damages), loss of revenue due to inadequate patent protection or inability to supply GSK products, and regulatory action such as fines, penalties, or loss of product authorisation. Any of these consequences could materially and adversely affect our financial results and cause loss of trust from our customers and patients.

Context

Research relating to animals can raise ethical concerns. While we attempt to address this proactively, animal studies remain a vital part of our research. In many cases, they are the only method that can be used to investigate the effects of a potential new medicine in a living body before it is tested in humans, and they are generally mandated by regulators and ethically imperative. Animal research can provide critical information about the causes of diseases and

how they develop. Nonetheless, we are continually seeking ways in which we can minimise our use of animals in research, whilst complying with regulatory requirements.

Clinical trials in healthy volunteers and patients are used to assess and demonstrate an investigational product's efficacy and safety or further evaluate the product once it has been approved for marketing. We also work with human biological samples. These samples are fundamental to the discovery, development and safety monitoring of our products.

The integrity of our data is essential to success in all stages of the research data lifecycle: design, generation, recording and management, analysis, reporting and storage and retrieval. Our research data is governed by legislation and regulatory requirements. Research data and supporting documents are core components at various stages of pipeline progression decision-making and form the content of regulatory submissions. Poor data integrity can compromise our research efforts and negatively impact company reputation.

There are innate complexities and interdependencies required for regulatory filings, particularly given our global research and development footprint. Continually changing and increasingly stringent submission requirements continue to increase the complexity of worldwide product registration.

Scientific engagement (SE), defined as the interaction and exchange of information between GSK and external communities to advance scientific and medical understanding, including the appropriate development and use of our products, is an essential part of scientific discourse. Such non-promotional engagement with external stakeholder groups is vital to GSK's mission and necessary for scientific and medical advance. SE activities are essential but present legal, regulatory, and reputational risk if the sharing of data, invited media coverage or payments to HCPs have, or are perceived to have, promotional intent.

A wide variety of biological materials are used by GSK in discovery, research and development phases. Through the Convention on Biological Diversity (CBD) and the Nagoya Protocol, the international community has established a global framework regulating access to, and use of, genetic resources of non-human origin in R&D. We support the principles of access and benefit sharing to genetic resources as outlined in the CBD and the Nagoya Protocol, recognising the importance of appropriate, effective and proportionate implementation measures at national and regional levels.

In addition, any loss of patent protection in a market for GSK's products developed through our R&D, including reducing the availability or scope of patent rights or compulsory licensing (in which a government forces a manufacturer to license its patents for specific products to a competitor), could materially and adversely affect our financial results in that market. Absence of adequate patent or data exclusivity protection, which could lead to, for example, competition from manufacturers of generic pharmaceutical products, could limit the opportunity to rely on such markets for future sales growth for our products, which could also materially and adversely impact our financial results. Following expiration of certain intellectual property rights, a generic manufacturer may lawfully produce a generic version of a product, and generic drug manufacturers have also exhibited a readiness to market generic versions of many of our most important products prior to the expiration of our patents. Introduction of generic products typically leads to a rapid and dramatic loss of sales and reduces our revenues and margins for our proprietary products. Moreover, in the US, it has become common for patent infringement actions to prompt claims that anti-trust laws have been violated during the prosecution of the patent or during litigation involving the defence of that patent.

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Third party oversight risk

Risk definition

Failure to maintain adequate governance and oversight over third party relationships and failure of third parties to meet their contractual, regulatory, confidentiality or other obligations.

Risk impact

Failure to adequately manage third party relationships could result in business disruption and exposure to risks ranging from sub-optimal contractual terms and conditions, to severe business and legal sanctions and/or significant reputational damage. Any of these consequences could materially and adversely affect our business operations and financial results.

Context

Third parties are critical to our business delivery and are an integral part of the solution to meeting our business objectives. We rely on third parties, including suppliers, advisors, distributors, individual contractors, licensees, and other pharmaceutical and biotechnology collaboration partners for discovery, manufacture, and marketing of our products and for supporting other important business processes.

These business relationships present a material risk. For example, we share critical and sensitive information such as marketing plans, clinical data, and employee data with specific third parties who are conducting the relevant outsourced business activities. Inadequate protection or misuse of this information by third parties could have significant business impact. Similarly, we use distributors and agents in a range of activities such as promotion and tendering which have inherent risks such as inappropriate promotion or corruption. Insufficient internal compliance and controls by the distributors could affect our reputation. These risks are further increased by the complexities of working with large numbers of third parties across a diverse geographical spread.

Environment, health and safety and sustainability

Risk definition

Failure to manage environment, health & safety and sustainability (EHS&S) risks in line with our objectives and policies and with relevant laws and regulations.

Risk impact

Failure to manage EHS&S risks could lead to significant harm to people, the environment and communities in which we operate, fines, failure to meet stakeholder expectations and regulatory requirements, litigation or regulatory action, and damage to the Group's reputation, which could materially and adversely affect our financial results.

Context

We are subject to health, safety and environmental laws of various jurisdictions. These laws impose duties to protect people, the environment, and the communities in which we operate, as well as potential obligations to remediate contaminated sites. We have also been identified as a potentially responsible party under the US Comprehensive Environmental Response Compensation and Liability Act at a number of sites for remediation costs relating to our use

or ownership of such sites in the US. Failure to manage these environmental risks properly could result in litigation, regulatory action and additional remedial costs that may materially and adversely affect our financial results. See Note 45 to the financial statements, *Legal proceedings*, for a discussion of the environmental related proceedings in which we are involved. We routinely accrue amounts related to our liabilities for such matters.

Information protection

Risk definition

The risk to GSK business activities if information becomes disclosed to those not authorised to see it, or if information or systems fail to be available or are corrupted, typically because of cybersecurity threats, although accident or malicious insider action may be contributory causes.

This also includes the risk of failure to collect, secure, and use personal information in accordance with data privacy laws.

Risk impact

Failure to adequately protect critical and sensitive systems and information may result in loss of commercial or strategic advantage and could materially affect our ongoing business operations, such as scientific research, clinical trials and manufacturing and supply chain activities. Failure to comply with data privacy laws could lead to adverse impact on individuals (for example financial loss, distress or prejudice). In both cases, damage to our reputation, litigation, or other business disruption including regulatory sanction could occur, which could materially and adversely affect our financial results.

Context

We rely on critical and sensitive systems and data, such as corporate strategic plans, intellectual property, manufacturing systems and trade secrets. There is the potential that our computer systems or information may be exposed to misuse or unauthorised disclosure.

We believe that the cyber security incidents that we have experienced to date have not resulted in significant disruptions to our operations, and have not had a significant adverse effect on our results of operations, or on third parties. However, as the threats evolve we cannot provide assurance that our significant efforts in protecting and monitoring our systems and information will always be successful in preventing compromise or disruption in future.

All parts of our business process personal information. The use of this information is critical to our operations and innovation, including the development and sale of our products, as well as management of our employees.

New and evolving laws and regulations, such as the European Union General Data Protection Regulation (GDPR), are likely to bring increased scrutiny of our data management.

Supply continuity and crisis management

Risk definition

Failure to deliver a continuous supply of compliant finished product; inability to respond effectively to a crisis incident in a timely manner to recover and sustain critical operations, including key supply chains.

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Risk impact

We recognise that failure to supply our products can adversely impact consumers and patients who rely on them. A material interruption of supply or exclusion from healthcare programmes could expose us to litigation or regulatory action and financial penalties that could adversely affect the Group's financial results. The Group's international operations, and those of its partners, expose our workforce, facilities, operations and information technology to potential disruption from natural events (e.g. storm or earthquake), man-made events (e.g. civil unrest, terrorism), and global emergencies (e.g. Ebola outbreak, Flu pandemic). It is important that we have robust crisis management and recovery plans in place to manage such events.

Context

Our supply chain operations are subject to review and approval by various regulatory agencies that effectively provide our licence to operate. Failure by our manufacturing and distribution facilities or by suppliers of key services and materials could lead to litigation or regulatory action such as product recalls and seizures, interruption of supply, delays in the approval of new products, and suspension of manufacturing operations pending resolution of manufacturing or logistics issues.

We rely on materials and services provided by third party suppliers to make our products, including active pharmaceutical ingredients (API), antigens, intermediates, commodities, and components for the manufacture and packaging of Pharmaceutical, Vaccine and Consumer Healthcare products. Some of the third party services procured, such as services provided by contract manufacturing and clinical research organisations to support development of key products, are important to ensure continuous operation of our businesses.

Although we undertake risk mitigation we recognise that certain events could nevertheless still result in delays or service interruptions. We use effective crisis management and business continuity planning to provide for the health and safety of our people and to minimise impact to us, by maintaining functional operations following a natural or man-made disaster, or a public health emergency.

Item 4. Information on the Company

4.A History and development of the company

The information set forth under the heading:

About GSK on the inside back cover;

Head Office and Registered Office on the outside back cover; and

Note 38 Acquisitions and disposals on pages 206 to 208 of the GSK Annual Report 2017 is incorporated herein by reference.

4.B Business overview

See Item 3.D Risk factors above;
In addition, the information set forth under the headings:

GSK at a glance on pages 2 to 3;

Chairman's statement on page 4;

CEO's statement on pages 5 to 7;

How we create long-term value on pages 8 to 9;

Industry trends on pages 10 to 11;

Our long-term priorities on pages 12 to 17;

Pharmaceuticals on pages 23 to 29;

Vaccines on pages 31 to 35;

Consumer Healthcare on pages 37 to 41;

Trust on pages 42 to 51;

Note 6 Segment information on pages 169 to 172;

Note 38 Acquisitions and disposals on pages 206 to 208;

Pharmaceutical products, competition and intellectual property on pages 254 to 255;

Vaccines products, competition and intellectual property on page 255; and

Consumer Healthcare products and competition on page 256
of the GSK Annual Report 2017 is incorporated herein by reference.

4.C Organizational structure
The information set forth under the heading:

Note 44 Principal Group companies on page 226; and

Group Companies on pages 276 to 286
of the GSK Annual Report 2017 is incorporated herein by reference.

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4.D Property, plant and equipment

The information set forth under the headings:

Property, plant and equipment within Group financial review on page 72;

Note 6 Segment information on page 171; and

Note 17 Property, plant and equipment on pages 181 to 182
of the GSK Annual Report 2017 is incorporated herein by reference.

Item 4A. **Unresolved Staff Comments**

Not applicable.

Item 5. **Operating and Financial Review and Prospects**

5.A Operating results

The information set forth under the headings:

Regulatory and political environment on page 11;

US tax reform on page 54;

Our approach to Brexit on page 55;

Non-controlling interests in ViiV Healthcare on page 59;

Cash generation and conversion on page 71;

Financial position and resources on pages 72 to 75;

Critical accounting policies on pages 76 to 77; and

Treasury policies on pages 77 to 78;
of the GSK Annual Report 2017 is incorporated herein by reference.

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The following tables reconcile Total results to Adjusted results. References in the GSK Annual Report 2017 to the reconciliations on page 67 of that report should be read to refer to the information in these tables.

Adjusted results reconciliation 31 December 2017

	Total results £m	Intangible asset amortisation £m	Intangible asset impairment £m	Major restructuring £m	Transaction -related £m	Divestments, significant legal and other items £m	US tax reform £m	Adjusted results (revised) £m
Gross profit	19,844	546	400	545	80			21,415
Operating profit	4,087	591	688	1,056	1,599	(119)	666	8,568
Profit before taxation	3,525	591	688	1,060	1,599	(205)	666	7,924
Profit after taxation	2,169	457	512	851	980	(456)	1,744	6,257
Earnings per share	31.4p	9.4p	10.5p	17.4p	19.2p	(9.4)p	33.3p	111.8p
Weighted average number of shares (millions)	4,886							4,886

The following adjustments are made in arriving at Adjusted gross profit

Cost of sales	(10,342)	546	400	545	80			(8,771)
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The following adjustments are made in arriving at Adjusted operating profit

Selling, general and administration	(9,672)			248		83		(9,341)
Research and development	(4,476)	45	288	263		18		(3,862)
Other operating income	(1,965)				1,519	(220)	666	

The following adjustments are made in arriving at Adjusted profit before tax

Net finance costs	(669)			4		8		(657)
Profit on disposal of associates	94					(94)		

The following adjustments are made in arriving at Adjusted profit after tax

Taxation	(1,356)	(134)	(176)	(209)	(619)	(251)	1,078	(1,667)
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Adjusted results reconciliation 31 December 2016

	Total results £m	Intangible asset amortisation £m	Intangible asset impairment £m	Major restructuring £m	Transaction -related £m	Divestments, significant legal and other items £m	Adjusted results (revised) £m
Gross profit	18,599	547	7	297	86	2	19,538
Operating profit	2,598	588	20	970	3,919	(424)	7,671
Profit before taxation	1,939	588	20	974	3,919	(416)	7,024
Profit after taxation	1,062	458	15	757	3,480	(246)	5,526
Earnings per share	18.8p	9.4p	0.3p	15.6p	61.6p	(5.1)p	100.6p

Weighted average number of shares (millions)	4,860						4,860
The following adjustments are made in arriving at Adjusted gross profit							
Cost of sales	(9,290)	547	7	297	86	2	(8,351)
The following adjustments are made in arriving at Adjusted operating profit							
Selling, general and administration	(9,366)			514		55	(8,797)
Research and development	(3,628)	41	13	159	(81)	28	(3,468)
Other operating income	(3,405)				3,914	(509)	
The following adjustments are made in arriving at Adjusted profit before tax							
Net finance costs	(664)			4		8	(652)
The following adjustments are made in arriving at Adjusted profit after tax							
Taxation	(877)	(130)	(5)	(217)	(439)	170	(1,498)

Table of Contents**Adjusted results reconciliation 31 December 2015**

	Total results £m	Intangible asset amortisation £m	Intangible asset impairment £m	Major restructuring £m	Transaction -related £m	Divestments, significant legal and other items £m	Adjusted results (revised) £m
Gross profit	15,070	522	147	563	89	12	16,403
Operating profit	10,322	563	206	1,891	2,238	(9,561)	5,659
Profit before taxation	10,526	563	206	1,896	2,238	(10,408)	5,021
Profit after taxation	8,372	402	156	1,455	1,886	(8,226)	4,045
Earnings per share	174.3p	8.3p	3.2p	30.1p	28.8p	(170.1)p	74.6p
Weighted average number of shares (millions)	4,831						4,831
The following adjustments are made in arriving at Adjusted gross profit							
Cost of sales	(8,853)	522	147	563	89	12	(7,520)
The following adjustments are made in arriving at Adjusted operating profit							
Selling, general and administration	(9,232)		7	1,009	88	151	(7,977)
Research and development	(3,560)	41	52	319		52	(3,096)
Other operating income	7,715				2,061	(9,776)	
The following adjustments are made in arriving at Adjusted profit before tax							
Net finance costs	(653)			5		12	(636)
Profit on disposal of associates	843					(843)	
Share of after tax profits/(losses) of associates and joint ventures	14					(16)	(2)
The following adjustments are made in arriving at Adjusted profit after tax							
Taxation	(2,154)	(161)	(50)	(441)	(352)	2,182	(976)

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Financial review 2017

The information set forth in the Group financial review on pages 52 to 78 of the GSK Annual Report 2017 is incorporated herein by reference excluding the following sections:

2018 guidance on page 55;

Our approach to tax on page 56;

Viability Statement on page 57;

Non-controlling interests in ViiV Healthcare on page 59;

Research and development under Total Results on page 65; and

Adjusting items on page 67.

2018 Guidance

We expect continued progress in 2018, including sales growth contributions from our new and recent product launches in HIV, Respiratory and Vaccines.

The expectation for 2018 Adjusted EPS growth is dependent on a number of factors including, in particular, uncertainties relating to the timing and extent of potential generic competition to Advair in the US.

In the event that no substitutable generic version of Advair is introduced to the US market in 2018, the Group expects 2018

Adjusted EPS growth of 4-7% at CER. This is based on an expected decline in 2018 in US Advair sales of 20-25%.

In the event of a mid-year introduction of a substitutable generic competitor to Advair in the US, the Group expects full-year 2018 US Advair sales of around £750 million at CER (US\$1.30/£1), with Adjusted EPS flat to down 3% at CER.

Both scenarios reflect the benefit of US tax reform with an expected 2018 effective tax rate on Adjusted profits of 19-20%. We are not able to give guidance for Total results as we cannot reliably forecast certain material elements of our Total results such as the future fair value movements on contingent consideration and put options, impairments of intangible assets and the future fair value movements on contingent consideration and put options arising from changes in foreign exchange rates, and therefore a reconciliation of the guidance for Adjusted results to equivalent guidance for Total results is not available without unreasonable effort.

Research and development

R&D expenditure was £4,476 million (14.8% of turnover), 23% higher at AER and 19% higher at CER than in 2016. This included charges of £106 million from the utilisation of the Priority Review Voucher in 2017 as well as increased investment in the progression of a number of mid and late-stage programmes. In addition, there were higher restructuring costs, primarily as a result of the provision for future clinical obligations as a result of the progressive withdrawal of Tanzeum and the decision to terminate the rights to sirukumab, and higher intangible asset impairments.

	2017 £m	2016 (revised) £m	£%	Growth CER%
Discovery	1,020	821	24	21
Development	1,450	1,249	16	13
Facilities and central support functions	536	558	(4)	(7)
Total Pharmaceuticals	3,006	2,628	14	11
Vaccines R&D	621	597	4	(2)
Consumer Healthcare R&D	235	243	(3)	(7)
Research and development	3,862	3,468	11	8
Items reconciling Total R&D to Adjusted R&D	614	160		
Total R&D	4,476	3,628		

The growth in Development expenditure was driven by the progression of a number of mid and late-stage programmes in HIV, Respiratory and Anaemia, together with the utilisation of the Priority Review Voucher in Q2 2017. The continuing high growth in Discovery expenditure reflected further investment in the early stage Oncology portfolio.

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Financial Review 2016

Reporting framework

Presentation of Group results

Our Group financial review discusses the operating and financial performance of the Group, cash flows and our financial position and resources. We compare the results for each year primarily with the results of the preceding year.

The geographic sales analysis has been revised to reflect a minor change to the Group's internal reporting structure made in 2017.

Total results

Total reported results represent the Group's overall performance. However, these results can contain material unusual or non-operational items that may obscure the key trends and factors determining the Group's operational performance. As a result, we also report Adjusted results, which is a non-IFRS measure.

Adjusted results

Core results have been renamed Adjusted results and, instead of all legal charges and expenses, only significant legal charges and expenses are excluded in order to present Adjusted results. All other legal charges and expenses are included in Adjusted results. Significant legal charges and expenses are those arising from the settlement of litigation or a government investigation that are not in the normal course and materially larger than more regularly occurring individual matters. They also include certain major legacy legal matters.

Adjusted results exclude the following items from Total results: amortisation and impairment of intangible assets (excluding computer software) and goodwill; major restructuring costs, including those costs following material acquisitions; significant legal charges (net of insurance recoveries) and expenses on the settlement of litigation and government investigations; transaction-related accounting adjustments for significant acquisitions, and other items, including disposals of associates, products and businesses, and other operating income other than royalty income, together with the tax effects of all of these items.

These items are excluded from Adjusted results either because their impact can be significant or because their exclusion improves comparabilities and consistency of reporting with the majority of our peer companies. This definition of Adjusted results aligns the Group's results better with the majority of our peer companies and how they report earnings.

Adjusted results reporting is utilised as one of the bases for internal performance reporting alongside Total results, cash flow generation and a number of other metrics. Adjusted results are presented and discussed in this Group financial review as we believe that Adjusted results are more representative of the performance of the Group's operations and allow the key trends and factors driving that performance to be more easily and clearly identified by shareholders. For the same reasons, the results of our four segments: Pharmaceuticals, Pharmaceuticals R&D, Vaccines and Consumer Healthcare are reported and measured on the same basis.

We also use a number of other adjusted, non-IFRS, measures to report the performance of our business. These measures are used by management for planning and reporting purposes and in discussions with and presentations to investment analysts and rating agencies and may not be directly comparable with similarly described measures used

by other companies. Non-IFRS measures may be considered in addition to, but not as a substitute for or superior to, information presented in accordance with IFRS.

CER growth

In order to illustrate underlying performance, it is our practice to discuss the results in terms of constant exchange rate (CER) growth. This represents growth calculated as if the exchange rates used to determine the results of overseas companies in Sterling had remained unchanged from those used in the previous year. CER% represents growth at constant exchange rates. £% or AER% represents growth at actual exchange rates.

Table of Contents**Group turnover**

Group turnover for the year increased 17% AER and 6% CER to £27,889 million, with Pharmaceuticals up 14% AER 3% CER, Vaccines up 26% AER 14% CER and Consumer Healthcare up 19% AER 9% CER, the growth in all three businesses still reflecting the impact of the Novartis transaction which completed on 2 March 2015. Sales of New Pharmaceutical and Vaccine products were £4,453 million, a Sterling increase of £2,465 million.

Group turnover by geographic region

	2016 (revised) £m	2015 (revised) £m	Growth £%	Growth CER%
US	10,197	8,222	24	10
Europe	7,476	6,435	16	6
International	10,216	9,266	10	1
	27,889	23,923	17	6

Group turnover outside of the US and Europe represented 37% of total Group turnover in 2016 (2015 39%).

Sales from new Pharmaceutical and Vaccine products

	2016 £m	2015 £m	Growth £ %	Growth CER%
Respiratory:				
<i>Relvar/Breo Ellipta</i>	620	257	>100	>100
<i>Anoro Ellipta</i>	201	79	>100	>100
<i>Arnuity Ellipta</i>	15	3	>100	>100
<i>Incruse Ellipta</i>	114	14	>100	>100
<i>Nucala</i>	102	1	>100	>100
CVMU:				
<i>Eperzan/Tanzeum</i>	121	41	>100	>100
HIV:				
<i>Tivicay</i>	953	588	62	45
<i>Triumeq</i>	1,735	730	>100	>100
Pharmaceuticals	3,861	1,713	>100	>100
<i>Bexsero</i>	390	115	>100	>100
<i>Menveo</i>	202	160	26	16
Vaccines	592	275	>100	96
	4,453	1,988	>100	>100

Sales of New Pharmaceutical and Vaccine products were £4,453 million and represented approximately 22% of Pharmaceuticals and Vaccines turnover.

Pharmaceuticals

Pharmaceuticals turnover was £16,104 million, up 14% AER and 3% CER. HIV sales grew 53% AER 37% CER. The Respiratory portfolio returned to growth with sales up 13% AER 2% CER, continuing the transition globally to newer products. Respiratory sales grew 20% AER 7% CER in the US and 16% AER 3% CER in International, but declined 2% AER 10% CER in Europe. Sales of New Pharmaceutical products were £3,861 million, a Sterling increase of £2,148 million, which more than offset the Sterling decline in *Seretide/Advair* sales of £196 million. Sales of Established products increased 1% AER but declined 8% CER, with declines in all regions, but particularly International, reflecting the loss of exclusivity for *Valtrex* in Canada, the impact of market reforms and the continued reshaping of the business in China and the impact of biennial price revisions in Japan. The overall impact of pricing to net sales of Pharmaceuticals was around -1%.

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US Pharmaceuticals turnover of £4,705 million grew 11% AER but declined 1% CER in 2016. This reflected a 20% AER 7% CER growth in the Respiratory portfolio, partly offset by the impact of generic competition to *Avodart*, down 58% AER 63% CER to £70 million, and *Lovaza*, down 54% AER 59% CER to £43 million. *Relenza* sales were also down 90% AER 91% CER to £7 million following a reallocation of government funding. Sales of new Respiratory products totalled £654 million and the growth of these products exceeded the decline in *Advair*. *Advair* sales fell 2% AER 13% CER to £1,829 million, representing a 7% volume decline and a 6% negative impact of price. *Ventolin* sales were up 38% AER 23% CER to £421 million, benefiting from competitor supply constraints early in the year, while *Flovent* sales were flat AER but declined 11% CER to £378 million, reflecting pricing pressures in the ICS market. *Benlysta* sales increased 33% AER 18% CER to £277 million with ongoing demand growth.

In Europe, Pharmaceuticals turnover increased 1% AER but declined 8% CER to £2,867 million. Respiratory sales declined 2% AER 10% CER to £1,383 million reflecting the ongoing transition to the new Respiratory portfolio and generic competition to *Seretide* which declined 18% AER 24% CER (16% volume decline and an 8% negative impact of price) to £835 million. This was partly offset by growth in the new Respiratory products, which recorded sales of £225 million. Established products sales were up 4% AER but down 4% CER to £513 million.

International Pharmaceuticals sales of £4,976 million were up 4% AER but down 5% CER. Sales in Emerging Markets grew 1% AER but declined 4% CER, impacted by the decline in the China business (down 4% AER 12% CER primarily as a result of the ongoing reshaping programme and broader Healthcare reforms including price reductions) but also by recent divestments in the International region, and the limitation of trading in Venezuela. In Japan, Pharmaceutical sales were up 17% AER but down 5% CER to £1,425 million, impacted by biennial price revisions on older products as well as supply interruptions to *Avodart* early in the year. Respiratory sales in Japan grew 27% AER 3% CER with strong growth of the new Respiratory products, up 100% AER 57% CER to £118 million, more than offsetting the decline in *Advair* sales.

Respiratory

Respiratory sales in 2016 increased 13% AER 2% CER to £6,510 million, reflecting the continuing transition of the Respiratory portfolio to newer products. Growth in the new Respiratory products, which recorded combined sales of £1,052 million, including *Relvar/Breo Ellipta* sales of £620 million, more than offset the decline in *Seretide/Advair*. *Flixotide/Flovent* sales grew 2% AER but decreased 8% CER to £637 million and *Ventolin* sales grew 27% AER 15% CER to £785 million.

In the US, Respiratory sales increased 20% AER 7% CER to £3,306 million (14% volume growth and a 7% negative impact of price). The growth of new Respiratory products more than offset the 2% AER 13% CER decline in *Advair* (7% volume decline and a 6% negative impact of price). The new *Ellipta* products recorded combined sales of £583 million, including *Breo Ellipta* sales of £344 million, with *Nucala*, the treatment for severe asthma, reporting sales of £71 million. Established Respiratory assets included *Ventolin*, with sales up 38% AER 23% CER to £421 million, and *Flovent*, which was flat AER but declined 11% CER to £378 million. *Ventolin* sales benefited from competitor supply constraints early in the year, while *Flovent* continued to be impacted by ongoing pricing pressures in the ICS market.

European Respiratory sales were down 2% AER 10% CER to £1,383 million, with *Seretide* sales down 18% AER 24% CER to £835 million (16% volume decline and an 8% negative impact of price), reflecting continued competition from generics and the transition of the Respiratory portfolio to newer products. The new Respiratory products recorded combined sales of £225 million in 2016, including *Relvar Ellipta* sales of £140 million.

Respiratory sales in the International region increased 16% AER 3% CER to £1,821 million with Emerging Markets up 13% AER 7% CER and Japan up 27% AER 3% CER. In Emerging Markets, sales of *Seretide* were up 3% AER but down 3% CER at £476 million, while *Ventolin* grew 20% AER 13% CER to £219 million. In Japan, *Adoair* grew 9% AER but declined 12% CER.

HIV

HIV sales increased 53% AER 37% CER to £3,556 million, with the US up 64% AER 46% CER, Europe up 42% AER 29% CER and International up 34% AER 21% CER. The growth in all three regions was driven by *Triumeq* and *Tivicay*.

Triumeq and *Tivicay* sales were £1,735 million and £953 million, respectively. *Epzicom/Kivexa* sales declined 19% AER 27% CER to £568 million, and *Selzentry* sales grew 1% AER but declined 9% CER to £125 million. There were also continued declines in the mature portfolio, mainly driven by generic competition to both *Combivir*, down 32% AER 38% CER to £23 million, and *Lexiva*, down 22% AER 26% CER to £51 million.

Immuno-inflammation

Immuno-inflammation sales grew 29% AER 15% CER to £340 million. Sales of *Benlysta* were £306 million, up 33% AER 19% CER, with sales in the US of £277 million, up 33% CER 18% AER.

Established products

Established products turnover grew 1% AER but fell 8% CER to £2,541 million, with *Valtrex* sales down 28% AER 37% CER to £118 million driven by a decline in Canada, down 91% AER 91% CER to £5 million, following the loss of exclusivity. *Zeffix* sales were down 17% AER 24% CER to £111 million and *Lovaza* sales in the US fell 54% AER 59% CER to £43 million.

The *Avodart* franchise was down 3% AER 14% CER to £635 million, primarily due to a 58% AER 63% CER decline in the US following the launch of generic competition in Q4 2015. Sales of *Eperzan/Tanzeum* were £121 million, primarily in the US. *Prolia* was divested at the end of 2015 and therefore no sales were recorded in 2016, compared with £43 million in 2015.

Dermatology sales declined 5% AER 12% CER to £393 million, adversely affected by supply constraints, while *Augmentin* sales grew 7% AER but were flat CER at £563 million. Sales of products for Rare diseases were up 14% AER but flat CER at £423 million, and included sales of *Volibris*, which were up 13% AER 1% CER to £172 million.

Vaccines

Vaccines sales grew 26% AER and 14% CER to £4,592 million. Growth benefited from the strong performance of *Bexsero* across all regions, higher demand for *Fluarix/FluLaval* in the US and International and a tender award for *Menveo* in International. Further growth was driven by *Synflorix* due to market expansion in International and a tender award in Europe. *Boostrix* sales benefited from higher demand in Europe and International. Growth was partly offset by *Infanrix/Pediarix* due to supply constraints in International, as well as unfavourable CDC stockpile movements for a number of products across the portfolio.

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In the US, sales grew by 27% AER 13% CER to £1,599 million. Growth was driven by market and share growth for *Bexsero*, *Menveo* and *Boostrix*, improved supply and higher demand for *Fluarix/FluLaval* and competitor supply issues that benefited *Infanrix/Pediarix*. This growth was partly offset by adverse stockpile movements on *Menveo* and an unfavourable comparison with the benefit to 2015 from CDC stockpile movements on *Infanrix/Pediarix*, *Boostrix* and *Rotarix*.

In Europe, sales grew 30% AER 18% CER to £1,423 million. Growth was driven primarily by *Bexsero* sales in private market channels in several countries including Spain and Italy, and in the UK following its inclusion in the NHS immunisation programme. *Boostrix* sales benefited from higher demand and competitor supply issues. Sales increased in Germany driven by improved supply of Hepatitis vaccines and higher demand for *Encepur* and *Rabipur*. Sales growth was also helped by a tender award for *Synflorix* in Poland but *Infanrix/Pediarix* sales were adversely impacted, mainly in Germany, France and Italy, by a competitor's return to the market during the year. Growth was also partly offset by the unfavourable comparison with 2015 when *Menveo* sales in the UK benefited from a catch-up tender win.

In International, sales grew 21% AER 10% CER to £1,570 million. Growth was driven primarily by *Synflorix*, due to market expansion in Nigeria, higher demand in Africa and private market demand in Asia. The growth in *Menveo* sales was driven by a tender award in Argentina and *Rotarix* sales benefited from higher demand in Brazil and Japan. Further growth in the region was driven by Brazil due to strong demand for *Bexsero*, *Menjugate*, and *Boostrix*. *Fluarix/FluLaval* sales grew due to higher uptake in Australia. Growth in the region was partly offset by lower sales of *Infanrix/Pediarix*, due to supply constraints, and lower Hepatitis vaccines sales, due to wholesaler destocking in China following the introduction of new private market distribution regulations.

Consumer Healthcare

The Consumer Healthcare business represents the Consumer Healthcare Joint Venture with Novartis together with the GSK Consumer Healthcare listed businesses in India and Nigeria, which are excluded from the Joint Venture. Results do not include the trading performance of the Nigeria beverages business in Q4 2016 following its sale on 30 September 2016.

Sales grew 19% AER and 9% CER to £7,193 million, benefiting significantly from the inclusion of sales of the former Novartis products for the first time for the first two months of the period. Strong performances were delivered by the power brands within the Oral health and Wellness categories and across all regions. Sales from innovation within the last three years represented approximately 13% of sales, with a particular contribution for *Flonase*, which was switched to OTC in Q1 2015. Other notable launches in 2016 included *Sensodyne True White* and *Excedrin Gel-tabs* in the US.

US sales grew 23% AER 9% CER to £1,761 million. *Sensodyne* delivered double-digit growth, benefiting from the launch in 2015 of *Repair and Protect* and the launch of *True White* in the first quarter of 2016, together with distribution gains for *Pronamel* and the newly launched *Pronamel Strong & Bright* variant. *Flonase OTC* delivered high single-digit growth, with a strong performance in the first half of 2016, driven by new formats, but impacted in the second half by increasing private label competition. *Excedrin* grew in double-digits, driven by the *Gel-tab* launch and new digital campaigns, and *Tums* also delivered double-digit growth, benefiting from supply improvements. This was partly offset by a decline in *Aquafresh* sales due to increased competitive pressures and a re-alignment of investment behind power brands.

Sales in Europe grew 22% AER 12% CER to £2,169 million, driven primarily by performances within the Wellness and Oral health categories. *Voltaren* continued to deliver double-digit growth at both AER and CER, driven largely by

the 12-hour variant and with strong performances across all key markets. Oral health sales grew in double digits AER and mid single-digits CER, with strong growth in *Sensodyne* and the Gum health portfolio, as well as 10% AER growth but a flat CER performance in *Aquafresh*, due to increased competitive pressures. At a market level, sales grew well in Italy, Scandinavia, the UK and Germany, partly offset by a decline in sales in CIS due to the impact on consumer spending of the weaker economic environment.

International sales of £3,263 million grew 16% AER 8% CER. Growth was delivered in many priority markets, primarily through the power brands across the Oral health and Wellness categories. This was partly offset by the impact of the sale of the Nigeria beverages business at the end of Q3 2016 as well as the affect of the restructuring of activity in Venezuela at the end of 2015. Growth of the International region was also affected by the combined impact on the Indian business of the demonetisation implemented in November and a more general slowing of the health food drink category which impacted the performance of the Nutrition category and *Horlicks* in particular. Elsewhere, strong growth was delivered in the Middle East, Latin America and China. The growth in the Middle East was driven by strong momentum across the power brands, particularly *Otrivin*, *Panadol* and *Sensodyne*. Double-digit performances were delivered in Brazil and Argentina as a result of better pricing and new product launches within Oral health. China delivered double-digit sales growth at AER and high single-digit growth at CER with contributions across the portfolio and with *Sensodyne* and *Voltaren* in particular benefiting from e-commerce and retail distribution expansion.

Total results

Cost of sales

Cost of sales as a percentage of turnover was 33.3%, down 3.7 percentage points in Sterling terms and 2.4 percentage points in CER terms compared with 2015. This reflected improved product mix, particularly the impact of higher HIV sales in Pharmaceuticals, but also in Vaccines and Consumer Healthcare and lower restructuring costs as well as an increased contribution from integration and restructuring savings in all three businesses.

These benefits were partly offset by continued adverse pricing pressure in Pharmaceuticals, primarily Respiratory, as well as continued investments in the supply chain.

Selling, general and administration

SG&A costs were 33.6% of turnover, 5.0 percentage points AER lower than in 2015 and 4.3 percentage points lower on a CER basis. This primarily reflected lower restructuring costs as well as the benefits from the Pharmaceuticals restructuring programme and integration benefits in Vaccines and Consumer Healthcare, partly offset by investment in promotional product support, particularly for new launches in Respiratory, HIV, Vaccines and Consumer Healthcare.

Table of Contents**Research and development**

R&D expenditure was £3,628 million (13% of turnover), 1.9% AER higher than in 2015 and 5.6% lower on a CER basis. This reflected the benefit from cost reduction programmes in Pharmaceuticals, Consumer Healthcare and Vaccines R&D and lower restructuring costs, partly offset by increased investment, particularly in Pharmaceuticals, reflecting investments in a number of new programmes and the costs of the acquired BMS HIV programme.

The operations of Pharmaceuticals R&D are broadly split into Discovery activities (up to the completion of phase IIa trials) and Development work (from phase IIb onwards) each supported by specific and common infrastructure and other shared services where appropriate. Phase IV costs and other administrative expenses are reported in SG&A and are not included in the table below.

	2016 £m	2015 £m	Growth £%	Growth CER%
Discovery	848	744	14	6
Development	1,275	1,136	12	4
Facilities and central support functions	505	433	17	9
Total Pharmaceuticals	2,628	2,313	14	5
Vaccines R&D	597	525	14	2
Consumer Healthcare R&D	243	258	(6)	(12)
Research and development	3,468	3,096	12	3
Items reconciling Total R&D to Adjusted R&D	160	464		
Research and development	3,628	3,560	2	(6)

The most significant factor driving Total Pharmaceuticals R&D growth was progression of the ViiV Healthcare HIV portfolio, including programmes acquired from BMS earlier in the year. The increase in Discovery was also driven by progression of the early stage Oncology portfolio and early investment in Bioelectronics. Development growth was primarily due to the start of new Phase III programmes, including HIV, respiratory and anaemia, partly offset by the benefit from R&D cost reduction programmes. The increase in facilities and central support functions costs partly reflected investment in new data warehousing and analytics to transform the way data is harnessed across R&D together with a re-allocation of central support costs.

Other operating income/(expense)

Net other operating expense of £3,405 million (2015 £7,715 million income) primarily reflected further accounting charges related to remeasurement of the contingent consideration liability related to the former Shionogi-ViiV Healthcare joint venture, along with remeasurement of the value attributable to the Consumer Healthcare Joint Venture put option and the liabilities first recognised in Q1 2016 for the Pfizer and Shionogi put options and preferential dividends in ViiV Healthcare. These remeasurements were driven by the unwinding of the discount applied to these future liabilities as well as updated trading forecasts and changes in the exchange rate assumptions used, updating them to period-end rates, which have increased the estimated total sterling values of GSK's Consumer Healthcare and ViiV Healthcare businesses.

These charges were partly offset by milestone income of £152 million in relation to the disposal of ofatumumab that was completed in 2015 and gains on a number of other divestments made during the year, including the remaining shares held by the Group in Aspen Pharmacare. The net other operating income of £7,715 million in 2015 included the profit on the disposal of the Oncology business to Novartis of £9,228 million.

Operating profit

Total operating profit was £2,598 million in 2016 compared with £10,322 million in 2015 which benefited from the net disposal gains recorded following the disposal of the Oncology business as part of the Novartis transaction.

Operating profit benefited from improved operating leverage driven by sales growth and a more favourable mix across all three businesses, together with lower levels of restructuring costs compared with 2015. However, there were further accounting charges related to remeasurement of the contingent consideration liability related to the former Shionogi-ViiV Healthcare joint venture, along with remeasurement of the value attributable to the Consumer Healthcare Joint Venture put option and the liabilities first recognised in Q1 2016 for the Pfizer and Shionogi put options and preferential dividends in ViiV Healthcare.

Contingent consideration cash payments are made to Shionogi and other companies, which reduce the balance sheet liability and hence are not recorded in the income statement. Total contingent consideration cash payments in 2016 amounted to £431 million (2015 £459 million). This included cash payments made by ViiV Healthcare to Shionogi in relation to its contingent consideration liability (including preferential dividends) which amounted to £417 million (2015 £159 million). In 2015 a milestone payment of £300 million was made to Novartis in relation to the Vaccines acquisition.

Table of Contents**Net finance costs**

	2016 £m	2015 £m
Finance income		
Interest and other income	70	99
Fair value movements	2	5
	72	104
Finance expense		
Interest expense	(701)	(719)
Unwinding of discounts on liabilities	(16)	(16)
Remeasurements and fair value movements	(4)	(8)
Other finance expense	(15)	(14)
	(736)	(757)

Share of after tax profits of associates and joint ventures

The share of profits of associates and joint ventures was £5 million (2015 £14 million).

Profit before taxation

Taking account of net finance costs and the share of profit of associates, profit before taxation was £1,939 million compared with £10,526 million in 2015.

Taxation

	2016 £m	2015 £m
UK current year charge	241	156
Rest of world current year charge	1,326	2,924
Charge in respect of prior periods	(149)	(508)
Total current taxation	1,418	2,572
Total deferred taxation	(541)	(418)
Taxation on Total profits	877	2,154

A tax charge of £877 million on total profit represented an effective tax rate of 45.2% (2015 20.5%) and reflected the non-deductibility of certain items included within the transaction-related adjustments, particularly the remeasurements of the put options related to ViiV Healthcare and the Consumer Healthcare Joint Venture.

Non-controlling interests

The allocation of earnings to non-controlling interests amounted to £150 million (2015 (£50) million), including the non-controlling interest allocations of Consumer Healthcare profits of £203 million (2015 £14 million) and the allocation of ViiV Healthcare losses of £83 million (2015 £143 million) including the impact of changes in the proportions of preferential dividends due to each shareholder based on the relative performance of different products in the year. The allocation also reflected the impact on the contribution of some of the Group's other entities with non-controlling interests primarily as a result of net losses in those entities arising from exchange.

Earnings per share

The Total earnings per share was 18.8p, compared with 174.3p in 2015. The decrease primarily reflected the benefit in 2015 from the disposal of the Oncology business to Novartis that closed in March 2015, together with the impact in 2016 of charges arising from increases in the valuations of the liabilities for contingent consideration and the put options associated with increases in the Sterling value of the Group's HIV and Consumer Healthcare businesses, partly offset by improved performance and reduced restructuring costs.

Dividends

The Board declared four interim dividends resulting in a total dividend for the year of 80 pence, in line with the dividend declared in 2015.

Items adjusted from Total results to present Adjusted results

Total results are adjusted for a number of items in order to present Adjusted results, as explained above. The items are discussed below.

Intangible asset amortisation and impairment

Intangible asset amortisation was £588 million, compared with £563 million in 2015. Intangible asset impairments of £20 million (2015 £206 million) included impairments of R&D and commercial assets. Both of these charges were non-cash items.

Table of Contents**Major restructuring and integration**

Major restructuring and integration charges of £970 million have been incurred (2015 £1,891 million), reflecting the phasing of planned restructuring projects following the completion of the Novartis transaction in 2015, as well as reduced charges for Pharmaceuticals restructuring projects as this programme enters its later stages. Cash payments made were £1,077 million (2015 £1,131 million) including the settlement of certain charges accrued in previous quarters.

Charges for the combined restructuring and integration programme to date are £3.7 billion, with cash charges of £2.9 billion and cash payments to date of £2.7 billion. The anticipated total cash charges of the combined programme were expected to be up to £3.65 billion and the non-cash charges up to £1.35 billion. The programme delivered incremental cost savings of £1.4 billion in 2016, including a currency benefit of £0.2 billion, and has now delivered approximately £3.0 billion of annual savings (including the currency benefit). The programme remains on track to deliver the originally targeted total annual savings during 2017. An estimated £300 million of additional cash charges are expected in 2017 along with some residual non-cash charges.

Transaction-related adjustments

Transaction-related adjustments resulted in a net charge of £3,919 million (2015 £2,238 million). This primarily reflected accounting charges for the remeasurement of the liability and the unwinding of the discounting effects on the value attributable to the Consumer Healthcare Joint Venture put option held by Novartis, the remeasurement and the unwinding of the discounting effects on the contingent consideration relating to the acquisition of the former Shionogi-ViiV Healthcare Joint Venture and the value attributable to the put options and preferential dividends payable to Pfizer and Shionogi.

Charge/(credit)	2016 £m	2015 £m
Consumer Healthcare Joint Venture put option	1,133	83
Contingent consideration on former Shionogi-ViiV Healthcare Joint Venture (including Shionogi preferential dividends)	2,162	1,874
ViiV Healthcare put options and Pfizer preferential dividends	577	
Other adjustments	47	281
Total transaction-related charges	3,919	2,238

The aggregate impact of unwinding the discount on these future and potential liabilities was £905 million (2015 £757 million), including £464 million on the Consumer Healthcare Joint Venture put option, £334 million on contingent consideration on the former Shionogi-ViiV Healthcare Joint Venture, and £58 million on the ViiV Healthcare put options and preference dividends. The remaining charge of £3,014 million was driven primarily by changes in exchange rate assumptions as well as updates to trading forecasts.

During 2016, GSK and Shionogi made several amendments to the Shareholders' Agreement for ViiV Healthcare regarding the Shionogi put option and the GSK call option. The estimated liability for Shionogi's put option was initially recognised on GSK's balance sheet at the end of Q1 2016 and de-recognised in December 2016, directly to equity, when it stood at £1,244 million.

Contingent consideration cash payments are made to Shionogi and other companies, which reduce the balance sheet liability and hence are not recorded in the income statement. Total contingent consideration cash payments in 2016 amounted to £431 million (2015 £459 million). This included cash payments made by ViiV Healthcare to Shionogi in relation to its contingent consideration liability (including preferential dividends) which amounted to £417 million (2015 £159 million). In 2015 a milestone payment of £300 million was made to Novartis.

Divestments, significant legal charges and other items

Divestments and other items included equity investment disposals, including the disposal of the remaining Aspen Pharmacare investment, dividends and impairments, milestone income on ofatumumab, a number of other asset disposals, and certain other adjusting items. Significant legal charges of £62 million (2015 £151 million) include the benefit of the settlement of existing matters as well as provisions for ongoing litigation. Significant legal cash payments were £102 million (2015 £285 million). Divestments and other items in 2015 included the profit on the disposal of the Oncology business to Novartis.

Adjusted results

We use Adjusted results, which is a non-IFRS measure, among other metrics including Total results and cash flow generation, to manage the performance of the Group. Non-IFRS measures may be considered in addition to, but not as a substitute for or superior to, information presented in accordance with IFRS. The definition of Adjusted results is set out above and reconciliations of Total results to Adjusted results are presented on pages 9 and 10.

Table of Contents**Cost of sales**

	2016		2015		Growth	Growth
	£m	% of turnover	£m	% of turnover	£%	CER%
Cost of sales	(8,351)	(29.9)	(7,520)	(31.4)	11	5

Cost of sales as a percentage of turnover was 29.9%, down 1.5 percentage points in Sterling terms and 0.3 percentage points in CER terms compared with 2015. This reflected improved product mix, particularly the impact of higher HIV sales in Pharmaceuticals, but also in Vaccines and Consumer Healthcare, as well as an increased contribution from integration and restructuring savings in all three businesses, partly offset by continued adverse pricing pressure in Pharmaceuticals, primarily Respiratory, as well as continued investments in the supply chain.

Selling, general and administration

	2016		2015		Growth	Growth
	(revised) £m	% of turnover	(revised) £m	% of turnover	£%	CER%
Selling, general and administration	(8,797)	(31.5)	(7,977)	(33.3)	10	2

SG&A costs were 31.5% of turnover, 1.8 percentage points lower in Sterling terms than in 2015 and 1.1 percentage points lower on a CER basis. This primarily reflected tight control of ongoing costs as well as the benefits from the Pharmaceuticals restructuring programme and integration benefits in Vaccines and Consumer Healthcare, partly offset by investment in promotional product support, particularly for new launches in Respiratory, HIV, Vaccines and Consumer Healthcare.

Research and development

	2016		2015		Growth	Growth
	£m	% of turnover	£m	% of turnover	£%	CER%
Research and development	(3,468)	(12.4)	(3,096)	(12.9)	12	3

R&D expenditure was £3,468 million (12.4% of turnover), 12% AER higher than in 2015 and 3% higher on a CER basis, reflecting increased investment, particularly in Total Pharmaceuticals. The operations of Pharmaceuticals R&D are broadly split into Discovery activities (up to the completion of phase IIa trials) and Development work (from phase IIb onwards) each supported by specific and common infrastructure and other shared services where appropriate. Phase IV costs and other administrative expenses are reported in SG&A and are not included in the table below.

	2016 £m	2015 £m	Growth £%	Growth CER%
Discovery	848	744	14	6
Development	1,275	1,136	12	4
Facilities and central support functions	505	433	17	9
Total Pharmaceuticals	2,628	2,313	14	5
Vaccines R&D	597	525	14	2
Consumer Healthcare R&D	243	258	(6)	(12)
Research and development	3,468	3,096	12	3

The most significant factor driving Total Pharmaceuticals R&D growth was progression of the ViiV Healthcare HIV portfolio, including programmes acquired from BMS earlier in the year. The increase in Discovery was also driven by progression of the early stage Oncology portfolio and early investment in Bioelectronics. Development growth was primarily due to the start of new Phase III programmes, including HIV, respiratory and anaemia, partly offset by the benefit from R&D cost reduction programmes. The increase in facilities and central support functions costs partly reflected investment in new data warehousing and analytics to transform the way data is harnessed across R&D together with a re-allocation of central support costs.

Royalty income

Royalty income was £398 million (2015 £329 million) primarily reflecting increased royalty income from Gardasil sales as well as the benefit of a catch-up adjustment to prior-year estimates.

Table of Contents**Adjusted operating profit**

Adjusted operating profit was £7,671 million, up 36% AER 14% CER on a turnover increase of 17% AER 6% CER. The Adjusted operating margin of 27.5% was 3.8 percentage points higher in Sterling terms than in 2015 and 1.8 percentage points higher on a CER basis. This reflected improved operating leverage driven by sales growth and a more favourable mix across all three businesses as well as delivery of restructuring and integration benefits and tight control of ongoing costs, partly offset by continued price pressure, particularly in Respiratory, and supply chain and R&D investments.

Adjusted operating profit by business

	2016		2015		Growth	Growth
	(revised)	Margin	(revised)	Margin		
	£m	%	£m	%	£%	CER%
Pharmaceuticals	7,976	49.5	6,449	45.6	24	7
Pharmaceuticals R&D	(2,488)		(2,168)		15	6
Pharmaceuticals	5,488	34.1	4,281	30.2	28	7
Vaccines	1,429	31.1	958	26.2	49	36
Consumer Healthcare	1,116	15.5	684	11.3	63	42
	8,033	28.8	5,923	24.8	36	17
Corporate & other unallocated costs	(362)		(264)		37	53
Adjusted operating profit	7,671	27.5	5,659	23.7	36	14

Pharmaceuticals

Pharmaceuticals operating profit was £5,488 million, 28% AER higher and 7% higher in CER terms than in 2015 on a turnover increase of 14% AER 3% CER. The operating margin of 34.1% was 3.9 percentage points higher in Sterling terms than in 2015 and 1.3 percentage points higher on a CER basis. This reflected a more favourable product mix, primarily driven by the growth in HIV sales, and the cost reduction benefit from the Pharmaceuticals restructuring programme, partly offset by increased investment in new product support, increased investment in R&D in a number of new programmes, the continued impact of lower prices, particularly in Respiratory, and the broader transition of the Respiratory portfolio.

Vaccines

Vaccines operating profit was £1,429 million, 49% AER higher and 36% higher than in 2015 in CER terms on a turnover increase of 26% AER 14% CER. The operating profit margin of 31.1% was 4.9 percentage points higher in Sterling terms than in 2015 and 5.3 percentage points higher on a CER basis. This reflected improved product mix and enhanced operating leverage from strong sales growth, together with restructuring and integration benefits in cost of sales, SG&A and R&D, and higher royalty income. These were partly offset by SG&A investments to support business growth, a number of inventory adjustments and additional supply chain investments.

Consumer Healthcare

Consumer Healthcare operating profit was £1,116 million, 63% AER higher and 42% higher than in 2015 in CER terms on a turnover increase of 19% AER 9% CER. The operating margin of 15.5% was 4.2 percentage points higher in Sterling terms than in 2015 and 3.4 percentage points higher on a CER basis. This reflected improvements in gross margin, reflecting mix benefits from the power brand strategy and better pricing, as well as a strong contribution from integration synergies benefiting both SG&A and R&D as a percentage of sales.

Net finance costs

	2016	2015
	£m	£m
Finance income		
Interest and other income	70	99
Fair value movements	2	5
	72	104
Finance expense		
Interest expense	(701)	(719)
Unwinding of discounts on liabilities	(4)	1
Remeasurements and fair value movements	(4)	(8)
Other finance expense	(15)	(14)
	(724)	(740)

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Net Adjusted finance expense was £652 million compared with £636 million in 2015, reflecting the translation effect of exchange rate movements on the reported Sterling costs of foreign currency denominated interest-bearing instruments.

Share of after tax profits/(losses) and joint ventures

The share of profits of associates and joint ventures was £5 million (2015 £2 million loss).

Adjusted profit before taxation

	2016		2015		Growth	Growth
	(revised)	% of	(revised)	% of	£%	CER%
	£m	turnover	£m	turnover		
Adjusted profit before tax	7,024	25.2	5,021	21.0	40	16

Taxation

Tax on Adjusted profit amounted to £1,498 million and represented an effective Adjusted tax rate of 21.3% (2015 19.4%). The increase in the effective rate primarily reflected the Group's changing earnings mix.

Non-controlling interests

The allocation of earnings to non-controlling interests amounted to £637 million (2015 £440 million), including the non-controlling interest allocations of Consumer Healthcare profits of £288 million (2015 £137 million) and the allocation of ViiV Healthcare profits, which increased to £324 million (2015 £224 million) including the impact of changes in the proportions of preferential dividends due to each shareholder based on the relative performance of different products in the year. The allocation also reflected the impact on the contribution of some of the Group's other entities with non-controlling interests primarily as a result of net losses in those entities arising from exchange.

Adjusted earnings per share

Adjusted EPS of 100.6p was up 35% at actual rates and 11% in CER terms compared with a 36% AER, 14% CER increase in operating profit, primarily reflecting the increased tax rate compared with 2015 and the greater contribution to growth from businesses in which there are significant non-controlling interests.

Financial position and resources**Property, plant and equipment**

Our business is science-based, technology-intensive and highly regulated by governmental authorities. We allocate significant financial resources to the renewal and maintenance of our property, plant and equipment to minimise risks of interruption to production and to ensure compliance with regulatory standards. A number of our processes use hazardous materials.

The total cost of our property, plant and equipment at 31 December 2016 was £22,164 million, with a net book value of £10,808 million. Of this, land and buildings represented £4,223 million, plant and equipment £3,481 million and

assets in construction £3,104 million. In 2016, we invested £1,544 million in new property, plant and equipment. This was mainly related to a large number of projects for the renewal, improvement and expansion of facilities at various worldwide sites. Property is mainly held freehold. New investment is financed from our liquid resources. At 31 December 2016, we had contractual commitments for future capital expenditure of £496 million and operating lease commitments of £840 million. We believe that our facilities are adequate for our current needs. We observe stringent procedures and use specialist skills to manage environmental risks from our activities.

Goodwill

Goodwill increased during the year to £5,965 million at 31 December 2016, from £5,162 million. The increase primarily reflected the impact of exchange movements.

Other intangible assets

Other intangible assets include the cost of intangibles acquired from third parties and computer software. The net book value of other intangible assets as at 31 December 2016 was £18,776 million (2015 £16,672 million). The increase in 2016 reflected the impact of exchange movements, development costs capitalised during the year of £240 million, partly offset by the amortisation and impairment of existing intangibles of £796 million and £29 million, respectively.

Investments in associates and joint ventures

We held investments in associates and joint ventures, with a carrying value at 31 December 2016 of £263 million (2015 £207 million). The market value at 31 December 2016 was £502 million (2015 £267 million). The largest of these investments was in Innoviva Inc. which had a book value at 31 December 2016 of £138 million (2015 £112 million). The market value at 31 December 2016 was £278 million.

Other investments

We held other investments with a carrying value at 31 December 2016 of £985 million (2015 £1,255 million). The decrease in the carrying value during the year was primarily due to the sale of the Group's remaining stake in Aspen Pharmacare Holdings Limited which had a book value at 31 December 2015 of £383 million. The most significant of the investments held at 31 December 2016 was in Theravance Biopharma, Inc. which had a book value at 31 December 2016 of £248 million (2015 £93 million). The other investments included equity stakes in companies with which we have research collaborations, which provide access to biotechnology developments of potential interest and interests in companies that arise from business divestments.

Derivative financial instruments: assets

We had current derivative financial instruments held at fair value of £156 million (2015 £125 million). The majority of this amount related to foreign exchange contracts both designated and not designated as accounting hedges.

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Inventories

Inventory of £5,102 million increased from £4,716 million in 2015, primarily reflected the impact of exchange movements.

Trade and other receivables

Trade and other receivables of £6,026 million increased from £5,615 million in 2015, primarily reflecting exchange movements.

Derivative financial instruments: liabilities

We held current derivative financial instruments at fair value of £194 million (2015 £153 million). This primarily related to foreign exchange contracts both designated and not designated as accounting hedges.

Trade and other payables

Trade and other payables were £11,964 million, up from £8,885 million in 2015, reflecting the Pfizer put option related to ViiV Healthcare recognised in the year, higher accruals for customer returns and rebates and the impact of exchange movements.

Provisions

We carried deferred tax provisions and other short-term and non-current provisions of £3,434 million at 31 December 2016

(2015 £3,286 million) of which £344 million (2015 £352 million) related to legal and other disputes and £554 million (2015 £816 million) related to the major restructuring programme. Provision has been made for legal and other disputes, indemnified disposal liabilities, employee related liabilities and the costs of the restructuring programme to the extent that at the balance sheet date a legal or constructive obligation existed and could be reliably estimated.

Pensions and other post-employment benefits

We account for pension and other post-employment arrangements in accordance with IAS 19. The deficits, net of surpluses before allowing for deferred taxation were £2,084 million (2015 £1,584 million) on pension arrangements and £1,693 million (2015 £1,387 million) on unfunded post-employment liabilities. The increases in the deficits were predominantly driven by lower discount rates that we used to discount the value of the liabilities, together with an increase in the UK inflation rate assumptions and a stronger US Dollar at the year end, partly offset by special funding contributions to the UK schemes and significant UK asset gains.

Other non-current liabilities

Other non-current liabilities of £8,445 million at 31 December 2016 (2015 £7,107 million) included £7,420 million (2015 £6,287 million) related to the present value of the estimated amount payable by us in the event of full exercise of Novartis' right to require us to acquire its 36.5% shareholding in the Consumer Healthcare Joint Venture.

Contingent consideration liabilities

Contingent consideration liabilities amounted to £5,896 million at 31 December 2016 (2015 £3,855 million), of which £5,304 million (2015 £3,409 million) represented the estimated present value of amounts payable to Shionogi relating to ViiV Healthcare and £545 million (2015 £405 million) represented the estimated present value of contingent consideration payable to Novartis related to the Vaccines acquisition. The liability due to Shionogi included £224 million in respect of preferential dividends of which £154 million was recognised directly in equity in the year. The liability for preferential dividends due to Pfizer at 31 December 2016 was £23 million.

Net debt

	2016 £m	2015 £m
Cash, cash equivalents and liquid investments	4,986	5,905
Borrowings repayable within one year	(4,129)	(1,308)
Borrowings repayable after one year	(14,661)	(15,324)
Net debt	(13,804)	(10,727)

At 31 December 2016, net debt was £13.8 billion, compared with £10.7 billion at 31 December 2015, comprising gross debt of £18.8 billion and cash and liquid investments of £5.0 billion. The increase in net debt primarily reflected a £2.2 billion adverse exchange impact from the translation of non-Sterling denominated debt and exchange on other financing items, dividends paid to shareholders of £4.9 billion including the special dividend of £1.0 billion, partly offset by free cash flow of £3.1 billion and asset disposals of £1.0 billion.

At 31 December 2016, our cash and liquid investments were held as follows:

	2016 £m	2015 £m
Bank balances and deposits	2,583	3,767
US Treasury and Treasury repo only money market funds	2,248	624
Liquidity funds	66	1,439
Government securities	89	75
	4,986	5,905

Cash and liquid investments of £3.2 billion (2015 £4.2 billion) were held centrally at 31 December 2016.

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5.B Liquidity and capital resources

The information set forth under the headings:

Cash generation and conversion on page 71;

Financial position and resources on pages 72 to 75; and

Treasury policies on pages 77 to 78
of the GSK Annual Report 2017 is incorporated herein by reference.

5.C Research and development, patents and licenses, etc.

The information set forth under Research and development under Financial Review 2017 in Item 5.A of this annual report on Form 20-F is incorporated herein by reference.

The information set forth under the headings:

Driving performance for profitable, sustainable growth within page 29;

Innovation within Pharmaceuticals on pages 24 to 27, Vaccines on pages 32 to 33 and Consumer Healthcare on pages 38 to 39;

Performance within Pharmaceuticals on pages 28 to 29; Vaccines on pages 34 to 35 and Consumer Healthcare on pages 40 to 41;

Research and development within page 65;

Pharmaceuticals and Vaccines product development pipeline on pages 251 to 253;

Pharmaceutical products, competition and intellectual property on pages 254 to 255;

Vaccines products, competition and intellectual property on page 255; and

Consumer Healthcare products and competition on page 256

of the GSK Annual Report 2017 is incorporated herein by reference.

5.D Trend information

The information set forth under the heading Financial Review 2017 in Item 5.A of this annual report on Form 20-F is incorporated herein by reference.

5.E Off-balance sheet arrangements

Not applicable.

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5.F Tabular disclosure of contractual obligations
The information set forth under the heading:

Contractual obligations and commitments on page 75
of the GSK Annual Report 2017 is incorporated herein by reference.

Item 6. **Directors, Senior Management and Employees**

6.A Directors and senior management
The information set forth under the headings:

Our Board on pages 82 to 85; and

Our Corporate Executive Team on pages 86 to 87
of the GSK Annual Report 2017 is incorporated herein by reference.

6.B Compensation

Remuneration report on pages 113 to 141; and

2017 Remuneration policy summary on pages 142 to 146
of the GSK Annual Report 2017 is incorporated herein by reference.

6.C Board practices
The information set forth under the heading:

Governance on pages 80 to 112; and

Additional remuneration disclosures on page 125; and

Donations to political organisations and political expenditure on page 275 of the GSK Annual Report 2017 is incorporated herein by reference.

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6.D Employees

The information set forth under the headings:

Engagement and Talent and Development on page 48;

Note 9 Employee costs on page 174;

Note 28 Pensions and other post-employment benefits on pages 190 to 197; and

Number of employees under Five year record on page 250
of the GSK Annual Report 2017 is incorporated herein by reference.

6.E Share ownership

The information set forth under the headings:

Note 43 Employee share schemes on pages 224 to 225;

Total remuneration for 2017 on pages 117 to 118;

Value earned from Long Term Incentives (LTIs) on page 122;

Update on performance of ongoing LTI awards on page 123; and

Directors interests in shares on pages 128 to 137
of the GSK Annual Report 2017 is incorporated herein by reference.

Item 7. Major Shareholders and Related Party Transactions

7.A Major shareholders

The information set forth under the headings:

Change of control and essential contracts on page 112;

Share capital and control on pages 267 to 268; and

Analysis of shareholdings at 31 December 2017 on page 268
of the GSK Annual Report 2017 is incorporated herein by reference.

7.B Related party transactions

The information set forth under the heading:

Note 35 Related party transactions on page 204
of the GSK Annual Report 2017 is incorporated herein by reference.

7.C Interests of experts and counsel

Not applicable.

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Item 8. Financial Information

8.A Consolidated Statements and Other Financial Information:
See item 18 below.

In addition, the information set forth under the headings:

 Note 45 Legal proceedings on pages 227 to 232; and

 Dividends on page 269
of the GSK Annual Report 2017 is incorporated herein by reference.

8.B Significant Changes
The information set forth under the heading Note 45 Legal proceedings on pages 227 to 232 of the GSK Annual Report 2017 is incorporated herein by reference.

Item 9. The Offer and Listing

9.A Offer and listing details
The information set forth under the headings:

 Market capitalisation on page 267;

 Share price on page 267; and

 Nature of trading market on page 268
of the GSK Annual Report 2017 is incorporated herein by reference.

9.B Plan of distribution
Not applicable.

9.C Markets

The information set forth under the headings:

Nature of trading market on page 268
of the GSK Annual Report 2017 is incorporated herein by reference.

9.D Selling shareholders

Not applicable.

9.E Dilution

Not applicable.

9.F Expenses of the issue

Not applicable.

Item 10. **Additional Information**

10.A Share Capital

Not applicable.

10.B Articles of Association of GlaxoSmithKline plc

The following is a summary of the principal provisions of the company's Articles of Association (the "Articles"). Shareholders should not rely on this summary, but should instead refer to the current Articles which are filed with the Registrar of Companies in the UK and can be viewed on the company's website. The Articles contain the fundamental provisions of the company's constitution, and the rules for the internal management and control of the company. The company has no statement of objects in its Articles and accordingly its objects are unrestricted in accordance with the provisions of the Companies Act 2006.

(a) Voting

All resolutions put to the vote at general meetings will be decided by poll. On a poll, every shareholder who is present in person or by proxy shall have one vote for every Ordinary Share of which he or she is the holder. In the case of

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joint holders of a share, the vote of the senior who tenders a vote, whether in person or by proxy, shall be accepted to the exclusion of the votes of the other joint holders, and seniority shall be determined by the order in which the names stand on the register. Unless the Directors otherwise decide, the right to attend a general meeting and voting rights may not be exercised by a shareholder who has not paid to the company all calls and other sums then payable by him or her in respect of his or her Ordinary Shares. The right to attend a general meeting and voting rights may not be exercised by a shareholder who is subject to an order under Section 794 of the Companies Act 2006 because he or she has failed to provide the company with information concerning his or her interests in Ordinary Shares within the prescribed period, as required by Section 793 of the Companies Act 2006.

(b) Transfer of Ordinary Shares

Any shareholder may transfer his or her Ordinary Shares which are in certificated form by an instrument of transfer in any usual form or in any other form which the Directors may approve. Such instrument must be properly signed and stamped or certified (or otherwise shown to the satisfaction of the Directors as being exempt from stamp duty) and lodged with the company together with the relevant share certificate(s) and such other evidence as the Directors may reasonably require to show the right of the transferor to make the transfer.

Any member may transfer title to his or her uncertificated Ordinary Shares by means of a relevant system, such as CREST.

The transferor of a share is deemed to remain the holder until the transferee's name is entered on the register. The Directors may decline to register any transfer of any Ordinary Share which is not fully paid.

Registration of a transfer of uncertificated Ordinary Shares may be refused in the circumstances set out in the uncertificated securities rules, and where, in the case of a transfer to joint holders, the number of joint holders to whom the uncertificated Ordinary Share is to be transferred exceeds four.

The Articles contain no other restrictions on the transfer of fully paid certificated Ordinary Shares provided: (i) the instrument of transfer is duly stamped or certified or otherwise shown to the satisfaction of the Directors to be exempt from stamp duty and is accompanied by the relevant share certificate and such other evidence of the right to transfer as the Directors may reasonably require; (ii) the transfer, if to joint transferees, is in favour of not more than four transferees; (iii) the instrument of transfer is in respect of only one class of shares; and (iv) the holder of the Ordinary Shares is not subject to an order under Section 794 of the Companies Act 2006. Notice of refusal to register a transfer must be sent to the transferee within two months of the instrument of transfer being lodged. The Directors may decline to register a transfer of Ordinary Shares by a person holding 0.25 per cent. or more of the existing Ordinary Shares if such person is subject to an order under Section 794 Companies Act 2006, after failure to provide the company with information concerning interests in those Ordinary Shares required to be provided under Section 793 of the Companies Act 2006, unless the transfer is carried out pursuant to an arm's length sale.

Provisions in the Articles will not apply to uncertificated Ordinary Shares to the extent that they are inconsistent with:

- (i) the holding of Ordinary Shares in uncertificated form;
- (ii) the transfer of title to Ordinary Shares by means of a system such as CREST; and

(iii) any provisions of the relevant regulations.

(c) Dividends and distribution of assets on liquidation

The profits of the company which are available for distribution and permitted by law to be distributed and which the company may by ordinary resolution from time to time declare, upon the recommendation of the Directors to distribute by way of dividend, in respect of any accounting reference period shall be distributed by way of dividend among holders of Ordinary Shares.

If in their opinion the company's financial position justifies such payments, the Directors may, as far as any applicable legislation allows, pay interim dividends on shares of any class of such amounts and in respect of such periods as they think fit. Except in so far as the rights attaching to, or the terms of issue of, any share otherwise provide, all dividends will be declared, apportioned and paid pro rata according to the amounts paid up on the shares during any portion of the period in respect of which the dividend is paid. As the company has only one class of Ordinary Shares, the holders of such Ordinary Shares will be entitled to participate in any surplus assets in a winding-up in proportion to their shareholdings.

(d) Variation of rights and changes in capital

Subject to the provisions of any statute (including any orders, regulations or other subordinate legislation made under it) from time to time in force concerning companies in so far as it applies to the company (the Companies Acts), the rights attached to any class of shares may be varied with the written consent of the holders of three-quarters in nominal value of the issued shares of that class (excluding any shares of that class held as treasury shares) or with the

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sanction of a special resolution passed at a separate meeting of the holders of shares of that class. At every such separate meeting, the provisions of the Articles relating to general meetings shall apply, except the necessary quorum shall be at least two persons entitled to vote and holding or representing as proxy at least one-third in nominal value of the issued shares of the relevant class(excluding any shares of that class held as treasury shares) (but provided that at any adjourned meeting one holder of shares of the relevant class present in person or by proxy shall be a quorum).

The rights conferred upon the holders of any Ordinary Shares shall not, unless otherwise expressly provided in the rights attaching to those Ordinary Shares, be deemed to be varied by the creation or issue of further shares ranking *pari passu* with them.

(e) Unclaimed dividends

All dividends or other sums payable on or in respect of any Ordinary Shares which remain unclaimed may be invested or otherwise made use of by the Directors for the benefit of the company until claimed. Unless the Directors decide otherwise, any dividend or other sums payable on or in respect of any Ordinary Shares unclaimed after a period of 12 years from the date when declared or became due for payment will be forfeited and revert to the company. The company may stop sending dividend cheques or warrants by post, or employ such other means of payment in respect of any Ordinary Shares, if at least two consecutive payments have remained uncashed or are returned undelivered or if one payment has remained uncashed or is returned undelivered and the company cannot establish a new address for the holder after making reasonable enquiries; however, in either case, the company must resume sending cheques or warrants or employ such other means of payment if the holder or any person entitled to the Ordinary Shares by transmission requests the resumption in writing.

(f) Untraced shareholders

The company may sell any certificated Shares in the company after advertising its intention in accordance with the requirements of the Articles and waiting for three months if the Ordinary Shares have been in issue for at least ten years and during that period at least three dividends have become payable on them and have not been claimed and, so far as any Director is aware, the company has not received any communication from the holder of the Ordinary Shares or any person entitled to them by transmission. Upon any such sale, the company will become indebted to the former holder of the Ordinary Shares or the person entitled to them by transmission for an amount equal to the net proceeds of sale unless forfeited. If no valid claim for the money has been received by the company during a period of six years from the date on which the relevant shares were sold by the company, the money will be forfeited and will belong to the company.

(g) Limitations on rights of non-resident or foreign shareholders

There are no limitations imposed by the Articles on the rights of non-resident or foreign shareholders except that there is no requirement for the company to serve notices on shareholders outside the United Kingdom and the United States, if no postal address in the United States or United Kingdom has been provided to the company.

(h) General meetings of shareholders

The Articles rely on the Companies Act 2006 provisions dealing with the calling of general meeting. The company is required by the Companies Act 2006 to hold an annual general meeting each year. General meetings of shareholders may be called as necessary by the Directors and must be called promptly upon receipt of a requisition from shareholders. Under the Companies Act 2006, an annual general meeting must be called by notice of at least 21 clear days. A general meeting other than an annual general meeting may be called on not less than 14 clear days notice

provided a special resolution reducing the notice period to 14 clear days has been passed at the immediately preceding annual general meeting or a general meeting held since that annual general meeting.

(i) Conflicts of interest

The Directors may, subject to the provisions of the Articles, authorise any matter which would otherwise involve a Director breaching his or her duty under the Companies Acts to avoid conflicts of interest (each a Conflict). A Director seeking authorisation in respect of a Conflict shall declare to the other Directors the nature and extent of his or her Conflict as soon as is reasonably practicable and shall provide the other Directors with such details of the matter as are necessary to decide how to address the Conflict. The board may resolve to authorise the relevant Director in relation to any matter the subject of a Conflict, save that the relevant Director and any other Director with a similar interest shall not count towards the quorum nor vote on any resolution giving such authority, and, if the other Directors so decide, shall be excluded from any meeting of the Directors while the Conflict is under consideration.

(j) Other Conflicts of Interest

Subject to the provisions of the Companies Acts, and provided the nature and extent of a Director's interest has been declared to the Directors, a Director may:

- (i) be party to, or otherwise interested in, any contract with the company, or in which the company has a direct or indirect interest;

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- (ii) hold any other office or place of profit with the company (except that of auditor) in conjunction with his office of director for such period and upon such terms, including remuneration, as the Directors may decide;
- (iii) act by himself or through a firm with which he is associated in a professional capacity for the company or any other company in which the company may be interested (otherwise than as auditor);
- (iv) be or become a director of, or employed by, or otherwise be interested in any holding company or subsidiary company of the company or any other company in which the company may be interested; and
- (v) be or become a director of any other company in which the company does not have an interest and which cannot reasonably be regarded as giving rise to a conflict of interest at the time of his appointment as director of that other company.

No contract in which a Director is interested shall be liable to be avoided, and any Director who is so interested is not liable to account to the company or its shareholders for any benefit realised by the contract by reason of the Director holding that office or of the fiduciary relationship thereby established. However, no Director may vote on, or be counted in the quorum, in relation to any resolution of the board relating specifically to his or her own appointment (including remuneration) or the terms of his or her termination of appointment or relating to any contract in which he or she has an interest (subject to certain exceptions).

Subject to the Companies Acts, the company may by ordinary resolution suspend or relax to any extent the provisions relating to directors' interests or restrictions on voting or ratify any transaction not duly authorised by reason of a contravention of such provisions.

(k) Directors' remuneration

Each of the Directors will be paid a fee at such rate as may from time to time be determined by the Directors, but the total fees paid to all of the directors for acting as directors (including amounts paid to any director who acts as chairman or is chairman of, or serves on any committee of the board of directors but excluding any amounts paid under any other provision of the Articles) shall not exceed the higher of:

- (i) £3 million a year; and
- (ii) any higher amount as the company may by ordinary resolution decide. Such fees may be satisfied in cash or in shares or any other non-cash form. Any Director who is appointed to any executive office, acts as Chairman, acts as senior independent director, acts as a scientific/medical expert on the board, is Chairman of, or serves on any committee of the Directors or performs any other services which the Directors consider to extend beyond the ordinary services of a Director shall be entitled to receive such remuneration (whether by way of salary, commission or otherwise) as the Directors may decide. Each Director may be paid reasonable travelling, hotel and other incidental

expenses he or she incurs in attending and returning from meetings of the Directors or committees of the Directors, or general meetings of the company, or otherwise incurred in connection with the performance of his or her duties for the company.

(l) Pensions and gratuities for Directors

The Directors or any committee authorised by the Directors may provide benefits by the payment of gratuities, pensions or insurance or in any other manner for any Director or former Director or their relations, connected persons or dependants, but no benefits (except those provided for by the Articles) may be granted to or in respect of a Director or former Director who has not been employed by or held an executive office or place of profit under the company or any of its subsidiary undertakings or their respective predecessors in business without the approval of an ordinary resolution of the company.

(m) Borrowing powers

Subject to the provisions of the Companies Act 2006, the Directors may exercise all the company's powers to borrow money; to mortgage or charge all or any of the company's undertaking, property (present and future), and uncalled capital; to issue debentures and other securities; and to give security either outright or as collateral security for any debt, liability or obligation of the company or of any third party.

(n) Retirement and removal of Directors

A Director is subject to re-election at every annual general meeting of the company if he or she:

- (i) held office at the time of the two previous annual general meetings and did not retire by rotation at either of them;

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- (ii) has held office, other than employment or executive office, for a continuous period of nine years or more; or

- (iii) he or she has been appointed by the Directors since the last annual general meeting.

In addition to any power of removal conferred by the Companies Acts the company may by special resolution remove any Director before the expiration of his or her period of office. No Director is required to retire by reason of his or her age, nor do any special formalities apply to the appointment or re-election of any Director who is over any age limit. No shareholding qualification for Directors shall be required.

(o) Vacation of office

The office of a director shall be vacated if:

- (i) he resigns or offers to resign, and the board resolves to accept such offer;
- (ii) his resignation is requested by all of the other directors and all of the other directors are not less than three in number;
- (iii) he is or has been suffering from mental or physical ill health and the board resolves that his office be vacated;
- (iv) he is absent without permission of the board from meetings of the board (whether or not an alternate director appointed by him attends) for six consecutive months and the board resolves that his office is vacated;
- (v) he becomes bankrupt or compounds with his creditors generally;
- (vi) he is prohibited by law from being a director; or
- (vii) he is removed from office pursuant to the Articles or the Companies Acts.

(p) Share rights

Subject to any rights attached to existing shares, shares may be issued with such rights and restrictions as the company may by ordinary resolution decide, or (if there is no such resolution or so far as it does not make specific provision) as the board may decide. Such rights and restrictions shall apply as if they were set out in the Articles. Redeemable shares may be issued, subject to any rights attached to existing shares. The board may determine the terms, conditions and manner of redemption of any redeemable share so issued. Such terms and conditions shall apply to the relevant shares as if they were set out in the Articles. Subject to the articles, any resolution passed by the shareholders and other shareholders rights, the Board may decide how to offer, allot, grant options over or otherwise deal with any

shares in the company.

10.C Material contracts

On April 22, 2014, GSK and Novartis AG (Novartis) entered into a three-part, inter-conditional transaction (the Transaction), pursuant to which they executed an implementation agreement (as subsequently amended, the Implementation Agreement), a contribution agreement relating to a consumer healthcare joint venture (as subsequently amended, the Contribution Agreement), a share and business sale agreement relating to the vaccines business of Novartis (as subsequently amended, the Vaccines SAPA), a sale and purchase agreement relating to the oncology business of GSK (as subsequently amended, the Oncology SAPA), a put option deed relating to the influenza vaccines business of Novartis (as subsequently amended, the Put Option Deed) and a shareholders agreement (the Shareholders Agreement, and, together with the Implementation Agreement, the Contribution Agreement, the Vaccines SAPA, the Oncology SAPA and the Put Option Deed, the Transaction Contracts).

Under the Vaccines SAPA, GSK purchased Novartis vaccines business (excluding Novartis influenza vaccines business). The purchase price for the business is up to US\$7,055,000,000 plus royalties. The US\$7,055,000,000 consists of US\$5,255,000,000 upfront and up to US\$1,800,000,000 in milestone payments.

Pursuant to the Shareholders Agreement entered into by GSK and Novartis at the closing of the Transaction, GSK has seven of eleven seats on Consumer Healthcare s board of directors, and Novartis has customary minority rights and exit rights at a pre-defined, market-based pricing mechanism.

GSK s shareholders approved the Transaction on December 18, 2014. The Transaction closed on March 2, 2015.

10.D Exchange controls

The information set forth under the heading:

Exchange controls and other limitations affecting security holders on page 267 of the GSK Annual Report 2017 is incorporated herein by reference.

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10.E Taxation

The information set forth under the heading:

Tax information for shareholders on pages 270 to 271
of the GSK Annual Report 2017 is incorporated herein by reference.

10.F Dividends and paying agents

Not applicable.

10.G Statement by experts

Not applicable.

10.H Documents on display

The information set forth under the heading:

Documents on display on page 270
of the GSK Annual Report 2017 is incorporated herein by reference.

10.I Subsidiary information

Not applicable.

Item 11. Quantitative and Qualitative Disclosures About Market Risk

The information set forth under the headings:

Treasury policies on pages 77 to 78; and

Note 42 Financial instruments and related disclosures on pages 213 to 223
of the GSK Annual Report 2017 is incorporated herein by reference.

Item 12. **Description of Securities Other than Equity Securities**

12.A Debt Securities

Not applicable.

12.B Warrants and Rights

Not applicable.

12.C Other Securities

Not applicable.

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12.D American Depositary Shares Fees and charges payable by ADR holders

The Bank of New York serves as the depositary (the "Depositary") for GSK's American Depositary Receipt ("ADR") programme. On April 6, 2015, GSK and the Depositary amended and restated the deposit agreement (the "Deposit Agreement") between GSK, the Depositary and owners and holders of ADRs. Pursuant to the Deposit Agreement, ADR holders may be required to pay various fees to the Depositary, and the Depositary may refuse to provide any service for which a fee is assessed until the applicable fee has been paid. In particular, the Depositary, under the terms of the Deposit Agreement, shall charge (i) a fee of \$5.00 or less per 100 American Depositary Shares (or portion thereof) for the delivery and surrender of American Depositary Shares, (ii) a fee of \$0.05 or less per American Depositary Share (or portion thereof) for any cash distribution made pursuant to this Deposit Agreement, (iii) a fee for the distribution of securities other than cash or shares and (iv) a fee of \$0.05 or less per American Depositary Share (or portion thereof) per annum for depositary services. In addition, the following charges shall be incurred by any party depositing or withdrawing Shares or surrendering ADRs or to whom American Depositary Shares are issued: (i) taxes and other governmental charges, (ii) such registration fees as may from time to time be in effect, (iii) certain cable, telex and facsimile transmission expenses, (iv) such expenses as are incurred by the Depositary in the conversion of foreign currency and (v) any other charges payable by the Depositary.

The Depositary may (i) withhold dividends or other distributions or sell any or all of the shares underlying the ADRs in order to satisfy any tax or governmental charge, (ii) deduct from any cash distribution any tax payable thereon or the cost of any currency conversion and (iii) collect any of its fees or charges by deduction from any cash distribution payable to ADR holders that are obligated to pay those fees or charges.

Direct and indirect payments by the Depositary

GSK receives payments from the Depositary in the form of (i) the reimbursement of expenses in connection with the administration, servicing and maintenance of the ADR programme, (ii) a portion of the fees collected by the Depositary for the issuance and cancellation of American Depositary Shares and (iii) a portion of any cash dividend fees and/or special dividend fees. In 2017, the Depositary made payments to GSK of approximately \$7.2 million, of which approximately \$6.1 million were related to expenses reimbursed and fees collected in connection with services provided in 2016.

Under certain circumstances, including removal of the Depositary or termination of the ADR programme by GSK, GSK is required to repay certain amounts paid to GSK and to compensate the Depositary for payments made or services provided on behalf of GSK.

PART II

Item 13. Defaults, Dividend Arrearages and Delinquencies Not applicable.

Item 14. Material Modifications to the Rights of Security Holders and Use of Proceeds Not applicable.

Item 15. Controls and Procedures

The information set forth under the heading:

Internal control framework on pages 105 to 106
of the GSK Annual Report 2017 is incorporated herein by reference.

US law and regulation

A number of provisions of US law and regulation apply to the company because our shares are quoted on the New York Stock Exchange (the NYSE) in the form of American Depositary Shares.

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NYSE rules

In general, the NYSE rules permit the company to follow UK corporate governance practices instead of those applied in the USA, provided that we explain any significant variations. This explanation is contained in Item 16.G of this Form 20-F. NYSE rules that came into effect in 2005 require us to file annual and interim written affirmations concerning the Audit & Risk Committee and our statement on significant differences in corporate governance.

Sarbanes-Oxley Act of 2002

Following a number of corporate and accounting scandals in the USA, Congress passed the Sarbanes-Oxley Act of 2002. Sarbanes-Oxley is a wide ranging piece of legislation concerned largely with financial reporting and corporate governance.

As recommended by the Securities and Exchange Commission (the SEC), the company has established a Disclosure Committee. The Committee reports to the CEO, the CFO and to the Audit & Risk Committee. It is chaired by the Company Secretary and the members consist of senior managers from finance, legal, corporate communications and investor relations.

External legal counsel, the external auditors and internal experts are invited to attend its meetings periodically. It has responsibility for considering the materiality of information and, on a timely basis, determining the disclosure of that information. It has responsibility for the timely filing of reports with the SEC and the formal review of the GSK Annual Report 2017 and Form 20-F. In 2017 the Committee met 18 times.

Sarbanes-Oxley requires that this annual report on Form 20-F contain a statement as to whether a member of our Audit & Risk Committee (ARC) is an audit committee financial expert as defined by Sarbanes-Oxley. For a summary regarding the Board's judgment on this matter, please refer to Item 16.A below and to page 85 under 'Judy Lewent, Skills and experience' and page 96 under 'Judy Lewent, Audit & Risk Committee Chair' of the GSK Annual Report 2017. Additional disclosure requirements arise under section 302 and section 404 of Sarbanes-Oxley in respect of disclosure controls and procedures and internal control over financial reporting.

Section 302: Corporate responsibility for financial reports

Sarbanes-Oxley also introduced a requirement for the CEO and the CFO to complete formal certifications, confirming that:

they have each reviewed the GSK Annual Report 2017 and Form 20-F;

based on their knowledge, the GSK Annual Report 2017 and Form 20-F contain no material misstatements or omissions;

based on their knowledge, the financial statements and other financial information fairly present, in all material respects, the financial condition, results of operations and cash flows as of the dates, and for the periods, presented in the GSK Annual Report 2017 and Form 20-F;

they are responsible for establishing and maintaining disclosure controls and procedures that ensure that material information is made known to them, and have evaluated the effectiveness of these controls and procedures as at the year-end, the results of such evaluation being contained in the GSK Annual Report 2017 and Form 20-F;

they are responsible for establishing and maintaining internal control over financial reporting that provides reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;

they have disclosed in the GSK Annual Report 2017 and Form 20-F any changes in internal controls over financial reporting during the period covered by the GSK Annual Report 2017 and Form 20-F that have materially affected, or are reasonably likely to affect materially, the company's internal control over financial reporting; and

they have disclosed, based on their most recent evaluation of internal control over financial reporting, to the external auditors and the ARC, all significant deficiencies and material weaknesses in the design or operation of internal controls over financial reporting which are reasonably likely to affect adversely the company's ability to record, process, summarise and report financial information, and any fraud (regardless of materiality) involving persons that have a significant role in the company's internal control over financial reporting.

The Group has carried out an evaluation under the supervision and with the participation of its management, including the CEO and CFO, of the effectiveness of the design and operation of the Group's disclosure controls and procedures as at 31 December 2017.

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There are inherent limitations to the effectiveness of any system of disclosure controls and procedures, including the possibility of human error and the circumvention or overriding of the controls and procedures. Accordingly, even effective disclosure controls and procedures can only provide reasonable assurance of achieving their control objectives.

Based on the Group's evaluation, the CEO and CFO have concluded that, as at December 31, 2017, the disclosure controls and procedures were effective to provide reasonable assurance that information required to be disclosed in the reports that the Group files and submits under the US Securities Exchange Act of 1934, as amended, is recorded, processed, summarised and reported as and when required and that it is accumulated and communicated to management, including the CEO and CFO, as appropriate, to allow timely decisions regarding disclosure.

The CEO and CFO completed these certifications on March 16, 2018.

Section 404: Management's annual report on internal control over financial reporting.

In accordance with the requirements of section 404 of Sarbanes-Oxley, the following report is provided by management in respect of the Company's internal control over financial reporting (as defined in Rules 13a-15(f) and 15d-15(f) under the US Securities Exchange Act of 1934):

management is responsible for establishing and maintaining adequate internal control over financial reporting for the Group. Internal control over financial reporting is designed to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with IFRS;

management conducted an evaluation of the effectiveness of internal control over financial reporting based on the framework in Internal Control – Integrated Framework (2013 Framework) issued by the Committee of Sponsoring Organizations of the Treadway Commission;

management has assessed the effectiveness of internal control over financial reporting, as at 31 December 2017 and has concluded that such internal control over financial reporting was effective. In addition, there have been no changes in the Group's internal control over financial reporting during 2017 that have materially affected, or are reasonably likely to affect materially, the Group's internal control over financial reporting; and

PricewaterhouseCoopers LLP, which has audited the consolidated financial statements of the Group for the year ended December 31, 2017, has also assessed the effectiveness of the Group's internal control over financial reporting under Auditing Standard No. 5 of the Public Company Accounting Oversight Board (United States). Their audit report can be found in Item 18 below.

Item 16.A Audit committee financial expert

The information set forth under the heading:

Membership , within the Audit & Risk Committee Report , on page 96; and

Sarbanes-Oxley Act of 2002 on page 274
of the GSK Annual Report 2017 is incorporated herein by reference.

Item 16.B Code of Ethics

The information set forth under the heading:

Code of Conduct and reporting lines on page 104
of the GSK Annual Report 2017 is incorporated herein by reference.

No waivers were granted from a provision of our code of ethics to an officer or person described in Item 16.B(a) that relates to one or more of the items set forth in Item 16.B(b) in 2017.

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Item 16.C Principal Accountant Fees and Services

16C.(a) Audit Fees

The information set forth in the table under the heading Fees payable to the company's auditor and its associates in the rows named Audit of parent company and consolidated financial statements, Audit of the company's subsidiaries and Attestation under s.404 of Sarbanes-Oxley Act 2002 on page 173 of the GSK Annual Report 2017 is incorporated herein by reference.

16C.(b) Audit-Related Fees

The information set forth in the table under the heading Fees payable to the company's auditor and its associates in the row named Other assurance services on page 173 of the GSK Annual Report 2017 is incorporated herein by reference. The other assurance services provided by the auditor relate to agreed upon procedures and other assurance services outside of statutory audit requirements.

16C.(c) Tax Fees

The information set forth in the table under the heading Fees payable to the company's auditor and its associates in the rows named Taxation compliance and Taxation advice on page 173 of the GSK Annual Report 2017 is incorporated herein by reference.

16C.(d) All Other Fees

The information set forth in the table under the heading Fees payable to the company's auditor and its associates in the row named All other services on page 173 of the GSK Annual Report 2017 is incorporated herein by reference. All other services provided by the auditor primarily related to advisory services for the year-ended 31 December 2017.

16C.(e) The information set forth under the heading Non-audit services on pages 102 to 103 of the GSK Annual Report 2017 is incorporated herein by reference.

16C.(f) Not applicable.

Item 16.D Exemptions from the Listing Standards for Audit Committees

Not applicable.

Item 16.E Purchases of Equity Securities by the Issuer and Affiliated Purchasers

Not applicable.

Item 16.F Change in Registrant's Certifying Accountant

GSK, through the Audit & Risk Committee, conducted an external audit tender in 2016 with a view to replacing PricewaterhouseCoopers LLP (PwC) from our 2018 financial year onwards. As disclosed in last year's Annual Report, PwC was not invited to participate in this audit tender process having regard to audit firm rotation requirements, as dictated by UK legislation. The audit tender process was completed in December 2016 when, following the recommendation of the Audit & Risk Committee, the Board announced that it would appoint Deloitte LLP (Deloitte) as GSK's new external auditor to undertake GSK's audit for the financial year ending 31 December 2018.

During the two years prior to 31 December 2017 and the subsequent interim period through 16 March 2018, (1) PwC has not issued any reports on the financial statements of the Company or the Group or on the effectiveness of internal control over financial reporting that contained an adverse opinion or a disclaimer of opinion, nor were the auditors reports of PwC qualified or modified as to uncertainty, audit scope, or accounting principles, and (2) there has not been any disagreement as that term is used in Item 16F(a)(1)(iv) of Form 20-F over any matter of accounting principles or practices, financial statement disclosure, or auditing scope or procedures, which disagreement if not resolved to PwC's satisfaction would have caused it to make reference to the subject matter of the disagreement in connection with its auditors' reports, or any reportable event as that term is used in Item 16F(a)(1)(v) of Form 20-F as described in the Group's Form 20-F during this two year period and through 16 March 2018

Further in the two years prior to 31 December 2017 and through 16 March 2018, GSK have not consulted with Deloitte regarding either: (i) the application of accounting principles to a specified transaction, either completed or proposed, or the type of audit opinion that might be rendered with respect to the consolidated financial statements of GSK; or (ii) any matter that was the subject of a disagreement as that term is used in Item 16F(a)(1)(iv) of Form 20-F or a reportable event as described in Item 16F(a)(1)(v) of Form 20-F.

Further information regarding external auditors' appointment is set forth under the headings 'External auditors' on page 96, 'Auditors' appointment' on pages 101 to 102 and 'Auditors' transition' on pages 103 to 104 of the GSK Annual Report 2017 and is incorporated herein by reference.

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PwC will resign after the firm has concluded the 2017 external audit process and the Audit & Risk Committee will recommend to the Board that Deloitte be appointed to fill the casual vacancy. GSK Shareholders will be invited to appoint Deloitte as GSK's new external auditors at the 2018 AGM to be held on 3 May 2018. Deloitte commenced transition activities, including observing PwC activity, as an independent audit firm on 4 July 2017.

GSK has provided PwC with a copy of the foregoing disclosure and has requested that PwC furnish GSK with a letter addressed to the SEC stating whether it agrees with such disclosure. A copy of the letter, dated 16 March 2018, is filed herewith as Exhibit 15.2.

Item 16.G Corporate Governance

Comparison of New York Stock Exchange Corporate Governance Standards and GlaxoSmithKline plc's corporate governance practice.

On November 4, 2003, the New York Stock Exchange (the "NYSE") adopted new corporate governance standards. The application of the NYSE's standards is restricted for foreign companies, recognizing that they have to comply with domestic requirements. As a foreign private issuer, GlaxoSmithKline plc ("GlaxoSmithKline" or the "Company") must comply with the following NYSE standards:

1. the Company must satisfy the audit committee requirements of the SEC;
2. the Chief Executive Officer (the "CEO") must promptly notify the NYSE in writing after any executive officer of the Company becomes aware of any non-compliance with any applicable provisions of the NYSE's corporate governance standards;
3. the Company must submit an annual affirmation to the NYSE affirming GlaxoSmithKline's compliance with applicable NYSE corporate governance standards, and submit interim affirmations to the NYSE notifying it of specified changes to the audit committee or a change to the status of the Company as a foreign private issuer; and

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4. the Company must provide a brief description of any significant differences between its corporate governance practices and those followed by US companies under the NYSE listing standards.

As a Company listed on the London Stock Exchange, GlaxoSmithKline is required to comply with the UK Listing Authority's Listing Rules (the Listing Rules) and to report non-compliance with the UK Corporate Governance Code (the UK Code).

The table below discloses differences between GlaxoSmithKline's current domestic corporate governance practices, which are based on the UK Code, and the NYSE corporate governance standards, applicable to US companies.

NYSE

Corporate Governance Standards

Director Independence (303A.01 of NYSE Manual)

1. Listed companies must have a majority of independent directors (as defined in Rule 10A-3 under the U.S. Securities Exchange Act of 1934, as amended (the Exchange Act)).

Description of differences between

GlaxoSmithKline's governance practice and the

NYSE Corporate Governance Standards

GlaxoSmithKline complies with the equivalent domestic requirements contained in the UK Code which was issued in April 2016.

The UK Code provides that the board of directors of GlaxoSmithKline (the Board) and its committees should have the appropriate balance of skills, experience, independence and

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NYSE Independence Tests (303A.02 of the NYSE Manual)

2. In order to tighten the definition of independent director for purposes of these standards:

(a) (i) No director qualifies as independent unless the board of directors affirmatively determines that the director has no material relationship with the listed company (either directly or as a partner, shareholder or officer of an organization that has a relationship with the company).

(ii) In addition, in affirmatively determining the independence of any director who will serve on the compensation committee of the listed company's board of directors, the board of directors must consider all factors specifically relevant to determining whether a director has a relationship to the listed company which is material to that director's ability to be independent from management in connection with the duties of a compensation committee member, including, but not limited to:

(A) the source of compensation of such director, including any consulting, advisory or other compensatory fee paid by the listed company to such director; and

(B) whether such director is affiliated with the listed company, a subsidiary of the listed company or an affiliate of a subsidiary of the listed company.

knowledge of the company to enable them to discharge their respective duties and responsibilities effectively (B.1). The Board should include an appropriate combination of Executive and Non-Executive Directors (and, in particular, independent Non-Executive Directors (for the purpose of the UK Code)) such that no individual or small group of individuals can dominate the Board's decision taking (B.1). At least half the Board, excluding the Chairman, should comprise Non-Executive Directors determined by the Board to be independent (B.1.2). The roles of Chairman and Chief Executive should not be exercised by the same individual. The division of responsibilities between the Chairman and Chief Executive should be clearly established, set out in writing and agreed by the Board (A.2.1).

The Board considers that Professor Sir Roy Anderson, Vindi Banga, Dr Vivienne Cox, Lynn Elsenhans, Dr Laurie Glimcher, Dr Jesse Goodman, Judy

Lewent, and Urs Rohner are independent for the purpose of the UK Code.

A majority of the Board members are independent Non-Executive Directors and, in accordance with the requirements of the UK Code, the Board has appointed one of the independent Non-Executive Directors as Senior Independent Director to provide a sounding board for the Chairman and act as an intermediary for other Directors where necessary (A.4.1). In January 2012 the Board adopted a formal written role specification for the Senior Independent Director.

GlaxoSmithKline complies with the corresponding domestic requirements contained in the UK Code, which sets out the principles for the Company to determine whether a director is independent.

The Board is required to determine and state its reasons for the determination of whether each Non-Executive Director is independent in character and judgment and whether there are relationships or circumstances which are likely to affect, or could appear to affect, the director's judgment. In undertaking this process, the Board is required, amongst other factors, to consider if the director:

- (a) has been an employee of GlaxoSmithKline within the last five years;
- (b) has, or has had within the last three years, a material business relationship with the Company either directly or as a partner, shareholder, director or senior employee of a body that has such a relationship with the Company;
- (c) has received or receives additional remuneration from the Company apart from a director's fee, participates in the Company's share option or a performance-related pay scheme, or is a member of the Company's pension scheme;
- (d) has close family ties with any of the Company's advisers, directors or senior employees;
- (e) holds cross-directorships or has significant links with other directors through involvement in other companies or bodies;
- (f) represents a significant shareholder; or
- (g) has served on the Board for more than nine years from the date of his or her first election, and is independent notwithstanding the existence of these relationships or circumstances (B.1.1).

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(b) In addition, a director is not independent if:

- (i) The director is, or has been within the last three years, an employee of the listed company, or an immediate family member is, or has been within the last three years, an executive officer, of the listed company.
- (ii) The director has received, or has an immediate family member who has received, during any twelve-month period within the last three years, more than \$120,000 in direct compensation from the listed company, other than director and committee fees and pension or other forms of deferred compensation for prior service (provided such compensation is not contingent in any way on continued service).
- (iii) (A) The director is a current partner or employee of a firm that is the listed company's internal or external auditor; (B) the director has an immediate family member who is a current partner of such a firm; (C) the director has an immediate family member who is a current employee of such a firm and personally works on the listed company's audit; or (D) the director or an immediate family member was within the last three years a partner or employee of such a firm and personally worked on the listed company's audit within that time.
- (iv) The director or an immediate family member is, or has been within the last three years, employed as an executive officer of another company where any of the listed company's present executive officers at the same time serves or served on that company's compensation committee.
- (v) The director is a current employee, or an immediate family member is a current executive officer, of a company that has made payments to, or received payments from, the listed company for property or services in an amount which, in any of the last three fiscal years, exceeds the greater of \$1 million, or 2% of such other company's consolidated gross revenues.

(For the purposes of these standards "executive officer" is defined to have the meaning specified for the term "officer" in Rule 16a-1(f) under the Securities Exchange Act of 1934, as amended, the "Exchange Act").

The Board considers all its Non-Executive Directors to be independent in character and judgment and has concluded that all its Non-Executive Directors are independent within the meaning of the UK Code. The Chairman satisfied the independence criteria on appointment in accordance with the UK Code (A.3.1).

GlaxoSmithKline complied with the UK Code requirement that all Directors should be subject to annual election or re-election by shareholders (B.7) at its Annual General Meeting in 2017, and intends to comply with this requirement at its 2018 Annual General Meeting.

The UK Code also provides that the Board should undertake a formal and rigorous annual evaluation of its own performance and that of its committees and individual Directors (B.6). Evaluation of the Board should consider the balance of skills, experience, independence and knowledge of the Company on the Board, its diversity, including gender, how the board works together as a unit, and other factors relevant to its effectiveness (B.6). GlaxoSmithKline has complied with this requirement. In addition, the evaluation of the Board should be externally facilitated at least

every three years and a statement should be made as to whether an external facilitator has any other connection with the Company and the external facilitator should be identified in the annual report (B.6.2).

Internally facilitated evaluations were conducted in 2015 and

2016. The Company conducted an externally facilitated evaluation in 2014 and 2017.

The UK Code provides that all Directors should receive an induction on joining the Board and should regularly update and refresh their skills and knowledge (B.4). The Chairman should ensure that new Directors receive a full, formal and tailored induction on joining the Board (B.4.1). The Chairman should regularly review and agree with each Director their training and development needs (B.4.2).

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Executive Sessions (303A.03 of the NYSE Manual)

3. To empower non-management directors to serve as a more effective check on management, the non-management directors of each listed company must meet at regularly scheduled executive sessions without management.

Nominating / Corporate Governance Committee (303A.04 of the NYSE Manual)

4. (a) Listed companies must have a nominating/corporate governance committee composed entirely of independent directors.
- (b) The nominating/corporate governance committee must have a written charter that addresses:
 - (i) the committee's purpose and responsibilities which, at minimum, must be to: identify individuals qualified to become board members, consistent with criteria approved by the board, and to select, or to recommend that the board select, the director nominees for the next annual meeting of shareholders; develop and recommend to the board a set of corporate governance guidelines applicable to the corporation; and oversee the evaluation of the board and management; and
 - (ii) an annual performance evaluation of the committee.

Meetings

GlaxoSmithKline complies with the equivalent domestic requirements set out in the UK Code, which requires that the Chairman of GlaxoSmithKline should hold meetings with the Non-Executive Directors without executives present. The Non-Executive Directors, led by the Senior Independent Director, also meet at least annually without the Chairman present to appraise the Chairman's performance (A.4.2).

The UK Code provides that the Chairman should promote a culture of openness and debate by facilitating the effective contribution of Non-Executive Directors, in particular, and ensuring constructive relations between Executive and Non-Executive Directors (A.3). In addition, the Chairman is responsible for ensuring that all Directors are made aware of their major shareholders' issues and concerns, and the Chairman should ensure that the views of the shareholders are communicated to the Board as a whole (E.1 and E.1.1).

Nominations Committee

GlaxoSmithKline complies with the corresponding domestic requirements set out in the UK Code, which requires that GlaxoSmithKline should have a Nominations Committee that is comprised of a majority of independent Non-Executive Directors (B.2.1).

GlaxoSmithKline's Nominations Committee has written terms of reference in accordance with the UK Code. The terms of reference are available on the Company's website and explain the Nominations Committee's role and the

authority delegated

to it by the Board (B.2.1). The Nominations Committee reviews the structure, size, diversity (including gender diversity), and

composition of the Board (evaluating the balance of skills, experience, independence and knowledge on the Board) and leads the process for the appointment of members to the Board and the Corporate Executive Team (the CET), and makes recommendations to the Board as appropriate. The Committee also monitors the planning of succession for the Board and Senior Management (B.2).

In compliance with the UK Code, the terms and conditions of appointment of Non-Executive Directors are available for inspection (B.3.2).

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Compensation Committee (303A.05 of the NYSE Manual)

5. (a) Listed companies must have a compensation committee composed entirely of independent directors. Compensation committee members must satisfy the additional independence requirements specific to compensation committee membership set forth in Section 2(a)(ii) in the Section titled Independence Tests above.
- (b) The compensation committee must have a written charter that addresses:
- (i) the committee's purpose and responsibilities which, at a minimum, must be to have direct responsibility to:
- (A) review and approve corporate goals and objectives relevant to CEO compensation, evaluate the CEO's performance in light of those goals and objectives, and, either as a committee or together with the other independent directors (as directed by the board), determine and approve the CEO's compensation level based on this evaluation;

The UK Code requires that a separate section in the Company's Annual Report describe the work of the Nominations Committee in discharging its duties, including the process it has used in relation to Board appointments (B.2.4). An explanation should be given if neither an external search consultancy nor open advertising has been used in the appointment of a chairman or a non-executive director. Where an external search consultancy has been used, it should be identified in the report and a statement should be made as to whether it has any other connection with the company (B.2.4). This section should include a description of the board's policy on diversity, including gender, any measurable objectives that it has set for implementing the policy, and progress on achieving the objectives (B.2.4). GlaxoSmithKline has complied with this requirement.

As described above, there is an annual Board evaluation exercise, which also includes evaluation of the Board's committees and individual Directors (B.6).

The Board is responsible for regularly reviewing its corporate governance standards and practices. The Company Secretary oversees corporate governance matters for the Group. The Company Secretary is responsible for advising the Board through the Chairman on all corporate governance matters. Domestic requirements do not mandate that GlaxoSmithKline establish a distinct corporate governance committee.

Remuneration Committee

GlaxoSmithKline complies with the equivalent domestic requirements set out in the UK Code, which requires that GlaxoSmithKline should have a Remuneration Committee that is comprised of at least three independent Non-Executive Directors (D.2.1).

GlaxoSmithKline's Remuneration Committee has written terms of reference in accordance with the UK Code, which explain the Remuneration's Committee's role and the authority delegated to it by the Board and which are available on

the Company's website (D.2.1). The Remuneration Committee determines the terms of service and remuneration of the Executive Directors and members of the CET and, with the assistance of external independent advisers, it evaluates and makes recommendations to the Board on overall executive remuneration policy (the Chairman and the CEO are responsible for evaluating and making recommendations to the Board on the remuneration of Non-Executive Directors). Where remuneration consultants are appointed, they should be

identified in the annual report and a statement should be made as to whether they have any other connection with the company

(D.2.1).

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(B) make recommendations to the board with respect to non-CEO executive officer compensation, and incentive-compensation and equity-based plans that are subject to board approval; and

(C) prepare the disclosure required by item 407(e)(5) or Regulation S-K under the Exchange Act;

(ii) an annual performance evaluation of the compensation committee.

(iii) The rights and responsibilities of the compensation committee set forth in Section 303A.05(c).

(c) (i) The compensation committee may, in its sole discretion, retain or obtain the advice of a compensation consultant, independent legal counsel or other adviser.

(ii) The compensation committee shall be directly responsible for the appointment, compensation and oversight of the work of any compensation consultant, independent legal counsel or other adviser retained by the compensation committee.

(iii) The listed company must provide for appropriate funding, as determined by the compensation committee, for payment of reasonable compensation to a compensation consultant, independent legal counsel or any other adviser retained by the compensation committee.

(iv) The compensation committee may select a compensation consultant, legal counsel or other adviser to the compensation committee only after taking into consideration, all factors relevant to that person's independence from management, including the following:

(A) The provision of other services to the listed company by the person that employs the compensation consultant, legal counsel or other adviser;

(B) The amount of fees received from the listed company by the person that employs the compensation consultant, legal counsel or other adviser, as a percentage of the total revenue of the person that employs the compensation consultant, legal counsel or other adviser;

The UK Code provides that the Remuneration Committee:

(a) should take care to recognise and manage conflicts of interest when receiving views from Executive Directors or senior management, or consulting the Chief Executive about its proposals (D.2) and should have delegated responsibility for setting remuneration for all Executive Directors and the Chairman, including pension rights and any compensation payments (D.2.2);

- (b) should recommend and monitor the level and structure of remuneration for senior management (D.2.2);
- (c) should consider what compensation commitments (including pension contributions and all other elements) the directors' terms of appointment would entail in the event of early termination (D.1.4.);
- (d) should invite shareholders specifically to approve all new long-term incentive schemes and significant changes to existing schemes (D.2.4.);
- (e) should judge where to position the Company relative to other companies and should be sensitive to pay and employment conditions elsewhere in the group, especially when determining annual salary increases (D.1); and
- (f) should consider whether the Directors should be eligible for annual bonuses and benefits under long-term incentive schemes and determine an appropriate balance between fixed and performance-related immediate and deferred remuneration bearing in mind that performance-related elements of Executive Directors' remuneration should be designed to promote the long-term success of the Company and be transparent, stretching and rigorously applied (D.1, D.1.1 and Schedule A). Incentive schemes should include provisions that would enable the Company to recover sums paid or withhold the payment of any sum, and specify the circumstances in which it would be appropriate to do so (D.1.1).

The UK Code requires that pay-outs under incentive schemes should be subject to relevant and stretching performance criteria, including non-financial performance criteria where appropriate and remuneration incentives should be compatible with the Company's risk policies and systems (Schedule A). In addition, remuneration of Non-Executive Directors should not include share options or other performance-related elements (D.1.3).

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- (C) The policies and procedures of the person that employs the compensation consultant, legal counsel or other adviser that are designed to prevent conflicts of interest;
- (D) Any business or personal relationship of the compensation consultant, legal counsel or other adviser with a member of the compensation committee;
- (E) Any stock of the listed company owned by the compensation consultant, legal counsel or other adviser; and
- (F) Any business or personal relationship of the compensation consultant, legal counsel, other adviser or the person employing the adviser with an executive officer of the listed company.

Audit Committee (303A.06 and 303A.07 of the NYSE Manual)

6. Listed companies must have an audit committee that satisfies the requirements of Rule 10A-3 under the Exchange Act.

As described above, there is an annual Board evaluation exercise, which also includes evaluation of the Board's committees (B.6).

Audit & Risk Committee

GlaxoSmithKline complies with equivalent domestic requirements set out in the UK Code, which requires that GlaxoSmithKline has an Audit & Risk Committee that is comprised of at least three independent Non-Executive Directors (C.3.1). The Company considers all members of the Audit & Risk Committee are independent. The Board has also satisfied itself, in line with the UK Code, that at least one member of the Audit & Risk Committee has recent and relevant financial experience and that the Audit & Risk Committee as a whole has competence relevant to the sector in which GlaxoSmithKline operates.

The UK Code requires the Audit & Risk Committee to:

- (a) monitor the integrity of the financial statements of the Company and any formal announcements relating to the Company's financial performance, reviewing significant financial reporting judgments contained in them (C.3.2);
- (b) review the Company's internal financial controls and internal control and risk management systems (C.3.2);
- (c) monitor and review the effectiveness of the Company's internal audit function (C.3.2);

- (d) Have primary responsibility for making a recommendation on the appointment, reappointment and removal of the external auditor (C.3.2);
- (e) make recommendations to the Board, for it to put to the shareholders for their approval in general meeting, in relation to the appointment, re-appointment and removal of the external auditor and to approve the remuneration and terms of engagement of the external auditor (C.3.2);

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- (f) review and monitor the external auditor's independence and objectivity and the effectiveness of the audit process, taking into consideration relevant UK professional and regulatory requirements (C.3.2);
 - (g) develop and implement policy on the engagement of external auditors to supply non-audit services, taking into account relevant ethical guidance regarding the provision of non-audit services by the external audit firm, and to report to the Board, identifying any matters in respect of which it considers that action or improvement is needed and making recommendations as to the steps to be taken (C.3.2);
 - (h) report to the Board on how it has discharged its responsibilities; and
 - (i) review arrangements by which the staff of the company may, in confidence, raise concerns about possible improprieties in matters of financial reporting or other matters (C.3.5).
- GlaxoSmithKline's Audit & Risk Committee meets the requirements of Rule 10A-3 in that:

each member of the Audit & Risk Committee is deemed to be independent in accordance with the Securities Exchange Act of 1934, as amended, and applicable NYSE and UK requirements;

the Audit & Risk Committee, amongst other things, is responsible for recommending the appointment, compensation, maintenance of independence and oversight of the work of any registered public accounting firm engaged for the purpose of preparing or issuing an audit report or performing other audit, review or attest services for the Company, and each such accounting firm must report directly to the Audit & Risk Committee;

the Audit & Risk Committee has established a procedure for the receipt, retention and treatment of complaints regarding accounting, internal accounting controls or auditing matters, and for the confidential, anonymous submission by employees of concerns regarding questionable accounting or auditing matters;

the Audit & Risk Committee has the authority to engage independent counsel and other advisors as it determines necessary to carry out its duties; and

GlaxoSmithKline must provide appropriate funding for the Audit & Risk Committee. The Board has determined that Judy Lewent has the appropriate qualifications and background to be an Audit Committee Financial Expert as defined in rules promulgated by the SEC under the Exchange Act.

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7. (a) The audit committee must have a minimum of three members. All audit committee members must satisfy the requirements for independence set out in Section 303A.02 and, in the absence of an applicable exemption, Rule 10A-3(b)(1) under the Exchange Act.

(b) The audit committee must have a written charter that addresses:

(i) the committee's purpose which, at minimum, must be to:

(A) assist board oversight of (1) the integrity of the listed company's financial statements, (2) the listed company's compliance with legal and regulatory requirements, (3) the independent auditor's qualifications and independence, and (4) the performance of the listed company's internal audit function and independent auditors (if the listed company does not yet have an internal audit function because it is availing itself of a transition period pursuant to Section 303A.00, the charter must provide that the committee will assist board oversight of the design and implementation of the internal audit function); and

(B) prepare disclosure regarding the audit committee's review and discussion of financial statements and certain other audit matters with management and auditors

(ii) the committee's responsibility to conduct an annual performance evaluation of the audit committee; and GlaxoSmithKline complies with the equivalent domestic requirements set out in the UK Code, which requires that the Audit & Risk Committee should be comprised of a minimum of three independent Non-Executive Directors (C.3.1).

GlaxoSmithKline's Audit & Risk Committee has written terms of reference in accordance with the UK Code. The terms of reference are available on the Company's website and explain the Audit & Risk Committee's role and the authority delegated to it by the Board (C.3.3). The Committee's main responsibilities include monitoring and reviewing the financial reporting process, the system of internal control and risk management, overseeing the identification and management of risks, the external and internal process and for monitoring compliance with laws, regulations and ethical codes of practice, including review throughout the year of integrated assurance reports comprising business unit and associated consolidated internal audit reports. Where requested by the board, the audit committee should provide advice on:

whether the annual report and accounts, taken as a whole, is fair, balanced and understandable and provides the information necessary for shareholders to assess the Company's performance, business model and strategy (C.3.4); and

when taking into account the Company's position and principal risks, how the prospects of the company have been assessed, over what period and why the period is regarded as appropriate. The Audit & Risk Committee should also advise whether there is a reasonable expectation that the company will be able to continue in operation and meet its liabilities when falling due over the said period, drawing attention to any

qualifications or assumptions as necessary prior to the directors making their statement in the annual report (C.2.2)

The UK Code requires that a separate section of the annual report should describe the work of the Committee in discharging its responsibilities (C.3.8).

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- (iii) the duties and responsibilities of the audit committee which, at a minimum, must include those set out in Rule 10A-3(b)(2), (3), (4) and (5) of the Exchange Act as well as to:
- (A) at least annually, obtain and review a report by the independent auditor describing: the firm's internal quality-control procedures; any material issues raised by the most recent internal quality-control review, or peer review, of the firm, or by any inquiry or investigation by governmental or professional authorities, within the preceding five years, respecting one or more independent audits carried out by the firm, and any steps taken to deal with any such issues; and (to assess the auditor's independence) all relationships between the independent auditor and the listed company;
 - (B) meet to review and discuss the listed company's annual audited financial statements and quarterly financial statements with management and the independent auditor, including reviewing the listed company's specific disclosures under Management's Discussion and Analysis of Financial Condition and Results of Operations ;
 - (C) discuss the listed company's earnings press releases, as well as financial information and earnings guidance provided to analysts and rating agencies;
 - (D) discuss policies with respect to risk assessment and risk management;
 - (E) meet separately, periodically, with management, with internal auditors (or other personnel responsible for the internal audit function) and with independent auditors;
 - (F) review with the independent auditor any audit problems or difficulties and management's response;
 - (G) set clear hiring policies for employees or former employees of the independent auditors; and
 - (H) report regularly to the board of directors.

(c) Each listed company must have an internal audit function.
The report should include:

the significant issues that the committee considered in relation to the financial statements, and how these issues were addressed (C.3.8);

an explanation of how it has assessed the effectiveness of the external audit process and the approach taken to the appointment or reappointment of the external auditor, information on the length of tenure of the current audit firm and when a tender was last conducted and advance notice of any retendering plans (C.3.8); and

if the external auditor provides non-audit services, an explanation of how auditor objectivity and independence are safeguarded (C.3.8).

Please see section 6 above for a description of the main role and responsibilities of the Audit & Risk Committee.

In accordance with the UK Code (C.3.6), the audit committee monitor and review the effectiveness of GlaxoSmithKline's internal audit function.

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Shareholder Approval of Equity Compensation Plans (303A.08 of the NYSE Manual)

8. Shareholders must be given the opportunity to vote on all equity-compensation plans and material revisions thereto, except for employment inducement awards, certain grants, plans and amendments in the context of mergers and acquisitions, and certain specific types of plans.

GlaxoSmithKline complies with corresponding domestic requirements in the Listing Rules, which mandate that the Company must seek shareholder approval for employee share schemes and significant changes to existing schemes, save in circumstances permitted by the Listing Rules (D.2.4 and Listing Rule 9.4). Please see section 5(d) above.

Corporate Governance Guidelines (303A.09 of the NYSE Manual)

9. Listed companies must adopt and disclose corporate governance guidelines.

GlaxoSmithKline complies with corresponding domestic requirements in the Listing Rules and the UK Code, which require that GlaxoSmithKline include an explanation in its Annual Report of how it complies with the principles of the UK Code and that it confirms that it complies with the UK Code's provisions or, where it does not, provide an explanation of how and why it does not comply (Listing Rule 9.8.6). In addition, GlaxoSmithKline is required to make certain mandatory corporate governance statements in the Directors' Report in accordance with the UK Listing Authority's Disclosure Guidance and Transparency Rules, DTR 7. With the exception of DTR 7.2.8AR and DTR 7.2.8BG (which apply to GlaxoSmithKline for the financial year beginning on 1 January 2017 and will be complied with in the 2017 Annual Report), GlaxoSmithKline has complied with these requirements in its 2016 Annual Report.

Code of Business Conduct and Ethics (303A.10 of the NYSE Manual)

10. Listed companies must adopt and disclose a code of business conduct and ethics for directors, officers and employees, and promptly disclose any waivers of the code for directors or executive officers.

Code of Conduct

GlaxoSmithKline's Code of Conduct for all employees, including the CEO, CFO and other senior financial officers, is available on the Company's website.

Foreign Private Issuer Disclosure (303A.11 of the NYSE Manual)

11. Listed foreign private issuers must disclose any significant ways in which their corporate governance practices differ from those followed by domestic companies under NYSE listing standards.

GlaxoSmithKline fulfils this requirement by publishing this document.

Listed foreign private issuers are required to provide this disclosure in the English language and in their

GlaxoSmithKline fulfils this requirement by including this disclosure in its annual report on Form 20-F.

annual reports filed on Form 20-F.

12. Certification Requirements (303A.12 of the NYSE Manual)

Each listed company and its CEO must file certain annual and interim certifications regarding compliance with the corporate governance requirements and certain other matters (although foreign private issuers are only required to comply with a subset of these requirements).

GlaxoSmithKline fulfils this requirement by filing the required certifications each year.

Item 16.H Mine Safety Disclosure

Not applicable.

PART III

Item 17 Financial Statements

Not applicable.

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Item 18 Financial Statements

The information set forth under the headings:

Consolidated income statement on page 158;

Consolidated statement of comprehensive income on page 158;

Consolidated balance sheet on page 159;

Consolidated statement of changes in equity on page 160;

Consolidated cash flow statement on page 161; and

Notes to the financial statements on pages 162 to 232
of the GSK Annual Report 2017 is incorporated herein by reference.

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Report on Form 20-F

Report of Independent Registered Public Accounting Firm

To the Board of Directors and Shareholders of GlaxoSmithKline plc

Opinions on the Financial Statements and Internal Control over Financial Reporting

We have audited the accompanying consolidated balance sheets of GlaxoSmithKline plc and its subsidiaries (the Company) at 31 December 2017 and 31 December 2016 and the related consolidated income statements, consolidated cash flow statements, consolidated statements of comprehensive income and consolidated statements of changes in equity for each of the three years in the period ended 31 December 2017, including the related notes, included in Exhibit 15.3 on pages 158 to 232 (collectively referred to as the consolidated financial statements). We have also audited the Company's internal control over financial reporting at 31 December 2017 based on criteria established in *Internal Control - Integrated Framework* (2013) issued by the Committee of Sponsoring Organizations of the Treadway Commission (COSO).

In our opinion, the consolidated financial statements referred to above present fairly, in all material respects, the financial position of the Company as of 31 December 2017 and 2016 and the results of its operations and its cash flows for each of the three years in the period ended 31 December 2017 in conformity with International Financial Reporting Standards as issued by the International Accounting Standards Board and in conformity with International Financial Reporting Standards as adopted by the European Union. Also in our opinion, the Company maintained, in all material respects, effective internal control over financial reporting at 31 December 2017 based on criteria established in *Internal Control - Integrated Framework* (2013) issued by the COSO.

Basis for Opinions

The Company's management is responsible for these consolidated financial statements, for maintaining effective internal control over financial reporting and for its assessment of the effectiveness of internal control over financial reporting, included in Management's annual report on internal control over financial reporting included in item 15 of 20-F. Our responsibility is to express opinions on the Company's consolidated financial statements and on the Company's internal control over financial reporting based on our audits. We are a public accounting firm registered with the Public Company Accounting Oversight Board (United States) (PCAOB) and are required to be independent with respect to the Company in accordance with the U.S. federal securities laws and the applicable rules and regulations of the Securities and Exchange Commission and the PCAOB.

We conducted our audits in accordance with the standards of the PCAOB. Those standards require that we plan and perform the audits to obtain reasonable assurance about whether the consolidated financial statements are free of material misstatement, whether due to error or fraud, and whether effective internal control over financial reporting was maintained in all material respects.

Our audits of the consolidated financial statements included performing procedures to assess the risks of material misstatement of the consolidated financial statements, whether due to error or fraud, and performing procedures that respond to those risks. Such procedures included examining, on a test basis, evidence regarding the amounts and

disclosures in the consolidated financial statements. Our audits also included evaluating the accounting principles used and significant estimates made by management, as well as evaluating the overall presentation of the consolidated financial statements. Our audit of internal control over financial reporting included obtaining an understanding of internal control over financial reporting, assessing the risk that a material weakness exists and testing and evaluating the design and operating effectiveness of internal control based on the assessed risk. Our audits also included performing such other procedures as we considered necessary in the circumstances. We believe that our audits provide a reasonable basis for our opinions.

Definition and Limitations of Internal Control over Financial Reporting

A company's internal control over financial reporting is a process designed to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles. A company's internal control over financial reporting includes those policies and procedures that (i) pertain to the maintenance of records that, in reasonable detail, accurately and fairly reflect the transactions and dispositions of the assets of the company; (ii) provide reasonable assurance that transactions are recorded as necessary to permit preparation of financial statements in accordance with generally accepted accounting principles and that receipts and expenditures of the company are being made only in accordance with authorisations of management and directors of the company; and (iii) provide reasonable assurance regarding prevention or timely detection of unauthorised acquisition, use, or disposition of the company's assets that could have a material effect on the financial statements.

Because of its inherent limitations, internal control over financial reporting may not prevent or detect misstatements. Also, projections of any evaluation of effectiveness to future periods are subject to the risk that controls may become inadequate because of changes in conditions or that the degree of compliance with the policies or procedures may deteriorate.

/s/ PricewaterhouseCoopers LLP

London, United Kingdom

16 March 2018

We have served as the Company or its merged predecessors' auditor since 1977. Since at least 1974, we also served as auditor of a company acquired by a merged predecessor of the Company.

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Item 19 Exhibits

- 1.1 Memorandum and Articles of Association of the Registrant as in effect on the date hereof.
- 2.1 Amended and Restated Deposit Agreement among the Registrant and The Bank of New York Mellon, as Depositary, and the owners and holders from time to time of the American Depositary Shares issued thereunder, including the form of American Depositary Receipt, is incorporated by reference to the post-effective amendment to the Registration Statement on Form F-6 (No. 333-148017) filed with the Commission on March 30, 2015.
- 4.1 UK Service Agreement between GlaxoSmithKline Services Unlimited and Simon Dingemans dated September 8, 2010 is incorporated by reference to Exhibit 4.7 to the Registrant's Annual Report on Form 20-F filed with the Commission on March 4, 2011.
- 4.2 UK Service Agreement between GlaxoSmithKline Services Unlimited and Emma N Walmsley dated December 20, 2016 is incorporated by reference to Exhibit 4.7 to the Registrant's Annual Report on Form 20-F filed with the Commission on March 17, 2017.
- 4.3 UK Service Agreement between GlaxoSmithKline Services Unlimited and Patrick John Thompson Vallance dated December 19, 2016 is incorporated by reference to Exhibit 4.8 to the Registrant's Annual Report on Form 20-F filed with the Commission on March 17, 2017.
- 4.4 UK Service Agreement between GlaxoSmithKline Services Unlimited and Emma N Walmsley dated March 29, 2017.
- 4.5 UK Service Agreement between GlaxoSmithKline LLC and Hal V. Barron dated December 16, 2017.
- 4.6 Share and Business Sale Agreement relating to the Vaccines Group made on April 22, 2014, as amended and restated on May 29, 2014, as amended on October 9, 2014, and as further amended and restated on March 1, 2015, between Novartis AG and GlaxoSmithKline plc is incorporated by reference to Exhibit 4.9 of the Registrant's Annual Report on Form 20-F filed with the Commission on March 18, 2016. Confidential portions of this exhibit have been omitted pursuant to a request for confidential treatment and filed separately with the SEC.
- 4.7 Shareholders' Agreement relating to GlaxoSmithKline Consumer Healthcare Holdings Limited made on March 2, 2015, among Setfirst Limited, Novartis Holding AG, Novartis Finance Corporation, GlaxoSmithKline plc, Novartis AG and GlaxoSmithKline Consumer Healthcare Holdings Limited is incorporated by reference to Exhibit 4.12 of the Registrant's Annual Report on Form 20-F filed with the Commission on March 18, 2016. Confidential portions of this exhibit have been omitted pursuant to a request for confidential treatment and filed separately with the SEC.
- 8.1 A list of the Registrant's principal subsidiaries is incorporated by reference to the information set forth under Group Companies 276 to 286 of the GSK Annual Report 2017 included as Exhibit 15.3.

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12.1	<u>Certification Required by Rule 13a-14(a) or 15d-14(a) under the Securities Exchange Act of 1934</u> Emma Walmsley
12.2	<u>Certification Required by Rule 13a-14(a) or 15d-14(a) under the Securities Exchange Act of 1934</u> Simon Dingemans.
13.1	<u>Certification Pursuant to Section 906 of the Sarbanes-Oxley Act of 2002 (Subsections (a) and (b) of Section 1350, Chapter 63 of Title 18, United States Code).</u>
15.1	<u>Consent of PricewaterhouseCoopers LLP.</u>
15.2	<u>Letter from PricewaterhouseCoopers LLP dated March 16, 2018.</u>
15.3*	<u>GSK Annual Report 2017.</u>
101.1**	101.1 Interactive Data Files (XBRL-Related Documents).

* Certain of the information included within Exhibit 15.3, which is provided pursuant to Rule 12b-23(a)(3) of the Securities Exchange Act of 1934, as amended, is incorporated by reference in this Form 20-F, as specified elsewhere in this Form 20-F. With the exception of the items and pages so specified, the GSK Annual Report 2017 is not deemed to be filed as part of this Form 20-F.

** As permitted by Rule 405(a)(2)(ii) of Regulation S-T, the registrant's XBRL (eXtensible Business Reporting Language) information will be furnished in an amendment to this Form 20-F that will be filed no more than 30 days after the date hereof. In accordance with Rule 402 of Regulation S-T, the information in these exhibits shall not be deemed to be filed for purposes of Section 18 of the Exchange Act, or otherwise subject to the liability of that section, and shall not be incorporated by reference into any registration statement or other document filed under the Securities Act, or the Exchange Act, except as shall be expressly set forth by specific reference in such filing.

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Signature

The registrant hereby certifies that it meets all of the requirements for filing on Form 20-F and that it has duly caused and authorized the undersigned to sign this Annual Report on its behalf.

GlaxoSmithKline plc

March 16, 2018

By: /s/ Simon Dingemans
Simon Dingemans
Chief Financial Officer