

Investor Information

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Financial record

Commercial Operations turnover by therapeutic area 2023

	Total			US			Europe			International		
	2023		Growth	2023		Growth	2023		Growth	2023		Growth
	£m	£%	CER%	£m	£%	CER%	£m	£%	CER%	£m	£%	CER%
Shingles	3,446	16	17	1,880	(4)	(4)	908	32	30	658	>100	>100
Shingrix	3,446	16	17	1,880	(4)	(4)	908	32	30	658	>100	>100
Meningitis	1,260	13	14	610	6	7	433	20	17	217	20	29
Bexsero	849	13	14	311	(7)	(6)	417	24	21	121	46	61
Menveo	380	10	12	299	25	25	12	(40)	(45)	69	(19)	(13)
Other	31	72	67	–	–	–	4	(20)	(20)	27	>100	>100
RSV	1,238	–	–	1,194	–	–	4	–	–	40	–	–
Arexvy	1,238	–	–	1,194	–	–	4	–	–	40	–	–
Influenza	504	(29)	(29)	371	(32)	(32)	39	(32)	(33)	94	(13)	(10)
Fluarix/FluLaval	504	(29)	(29)	371	(32)	(32)	39	(32)	(33)	94	(13)	(10)
Established Vaccines	3,266	6	7	1,254	8	9	742	3	2	1,270	5	7
Infanrix, Pediarix	554	(7)	(6)	291	(11)	(11)	121	(8)	(8)	142	4	10
Boostrix	614	3	4	394	9	10	122	(12)	(13)	98	2	4
Hepatitis	611	7	8	336	(2)	(1)	177	25	23	98	14	17
Rotarix	614	17	18	192	>100	>100	118	(3)	(5)	304	(2)	2
Synflorix	275	(10)	(10)	–	–	–	36	6	3	239	(12)	(12)
Priorix, Priorix Tetra, Varilrix	265	41	41	16	60	60	129	33	30	120	48	53
Cervarix	120	3	5	–	–	–	33	50	45	87	(8)	(4)
Others	213	13	11	25	14	9	6	(82)	(76)	182	37	34
Vaccines ex COVID	9,714	23	24	5,309	25	26	2,126	16	15	2,279	26	31
Pandemic vaccines	150	>100	>100	–	–	–	130	>100	>100	20	>100	>100
Pandemic adjuvant	150	>100	>100	–	–	–	130	>100	>100	20	>100	>100
Vaccines	9,864	24	25	5,309	25	26	2,256	20	18	2,299	27	31
HIV	6,444	12	13	4,283	14	14	1,423	9	7	738	8	16
Dolutegravir products	5,408	4	5	3,418	3	4	1,290	4	3	700	9	17
Tivicay	1,386	–	2	801	(3)	(2)	267	(2)	(4)	318	12	21
Triumeq	1,542	(14)	(14)	1,074	(12)	(11)	280	(22)	(24)	188	(15)	(11)
Juluca	661	4	4	511	3	4	136	7	6	14	(7)	(7)
Dovato	1,819	32	33	1,032	33	33	607	27	25	180	50	59
Rukobia	117	43	44	110	39	41	7	>100	>100	–	–	–
Cabenuva	708	>100	>100	587	100	>100	103	>100	>100	18	>100	>100
Apretude	149	>100	>100	149	>100	>100	–	–	–	–	–	–
Others	62	(35)	(33)	19	(39)	(42)	23	(18)	(25)	20	(44)	(31)
Respiratory/Immunology and Other	3,025	16	18	2,100	15	15	468	28	26	457	11	21
Nucala	1,655	16	18	978	11	11	383	28	26	294	21	33
Benlysta	1,349	18	19	1,121	18	19	99	19	18	129	13	25
Other	21	(48)	(42)	1	–	–	(14)	18	12	34	(40)	(33)
Oncology	731	21	23	396	27	27	289	14	13	46	28	61
Zejala	523	13	15	257	9	10	222	14	12	44	29	65
Blenrep	36	(69)	(69)	(2)	>(100)	>(100)	38	(27)	(27)	–	–	–
Jemperli	141	>100	>100	108	>100	>100	31	>100	>100	2	>100	>100
Ojjaara	33	–	–	33	–	–	–	–	–	–	–	–
Other	(2)	>(100)	>(100)	–	–	–	(2)	(100)	–	–	>(100)	(100)
Specialty Medicines ex COVID	10,200	14	15	6,779	15	15	2,180	13	11	1,241	10	19
Pandemic	44	(98)	(98)	10	(99)	(99)	3	(99)	(99)	31	(97)	(97)
Xevudy	44	(98)	(98)	10	(99)	(99)	3	(99)	(99)	31	(97)	(97)
Specialty Medicines	10,244	(9)	(8)	6,789	1	1	2,183	(8)	(10)	1,272	(41)	(36)
Respiratory	6,825	4	6	3,442	7	8	1,402	1	–	1,981	1	9
Arnuita Ellipta	36	(36)	(34)	29	(40)	(40)	–	–	–	7	(13)	–
Anoro Ellipta	557	15	16	269	15	16	193	17	15	95	12	20
Avamys/Veramyst	299	(7)	(4)	–	–	–	57	(12)	(14)	242	(5)	(2)
Flixotide/Flovent	451	(17)	(16)	283	(20)	(20)	70	(5)	(5)	98	(17)	(11)
Incruse Ellipta	162	(17)	(17)	78	(25)	(24)	59	(8)	(9)	25	(11)	(7)
Relvar/Breo Ellipta	1,103	(4)	(2)	436	(12)	(12)	366	5	4	301	–	8
Seretide/Advair	1,139	(2)	1	341	11	11	256	(11)	(12)	542	(4)	3
Trelegy Ellipta	2,202	27	29	1,606	28	29	275	17	16	321	34	44
Ventolin	749	(3)	–	400	(3)	(2)	100	(14)	(16)	249	2	11
Other Respiratory	127	(11)	(5)	–	(100)	(100)	26	(13)	(17)	101	(10)	(1)
Other General Medicines	3,395	(5)	2	280	(23)	(22)	723	4	2	2,392	(5)	6
Dermatology	363	(3)	4	–	–	–	107	–	(1)	256	(5)	6
Augmentin	628	9	17	–	–	–	186	23	21	442	4	16
Avodart	345	5	7	–	–	–	109	2	(1)	236	6	10
Lamictal	435	(15)	(13)	194	(27)	(27)	111	2	1	130	(5)	4
Other	1,624	(9)	1	86	(13)	(11)	210	(5)	(7)	1,328	(9)	3
General Medicines	10,220	1	5	3,722	4	5	2,125	2	1	4,373	(2)	7
Total Commercial Operations	30,328	3	5	15,820	9	9	6,564	3	2	7,944	(6)	1

Financial record continued

Commercial Operations turnover by therapeutic area 2022

	Total			US			Europe			International		
	2022		Growth	2022		Growth	2022		Growth	2022		Growth
	£m	£%	CER%	£m	£%	CER%	£m	£%	CER%	£m	£%	CER%
Shingles	2,958	72	60	1,964	46	32	688	>100	>100	306	>100	>100
<i>Shingrix</i>	2,958	72	60	1,964	46	32	688	>100	>100	306	>100	>100
Meningitis	1,116	16	11	573	26	14	362	2	3	181	18	20
<i>Bexsero</i>	753	16	12	333	32	19	337	3	4	83	20	23
<i>Menveo</i>	345	27	18	240	20	8	20	(5)	(10)	85	67	71
<i>Other</i>	18	(54)	(54)	–	–	–	5	–	–	13	(62)	(62)
Influenza	714	5	(4)	549	20	9	57	(44)	(44)	108	(11)	(16)
<i>Fluarix/Flulaval</i>	714	5	(4)	549	20	9	57	(44)	(44)	108	(11)	(16)
Established vaccines	3,085	4	–	1,157	18	7	720	3	4	1,208	(7)	8
<i>Infanrix, Pediarix</i>	594	9	3	327	8	(3)	131	13	13	136	10	6
<i>Boostrix</i>	594	14	7	360	33	20	138	(1)	(1)	96	(14)	(15)
<i>Hepatitis</i>	571	24	16	343	28	15	142	30	31	86	5	(1)
<i>Rotarix</i>	527	(3)	(3)	95	(14)	(23)	122	3	5	310	(1)	1
<i>Synflorix</i>	305	(15)	(15)	–	–	–	34	(24)	(22)	271	(13)	(14)
<i>Priorix, Priorix Tetra, Varilrix</i>	188	(28)	(29)	10	–	–	97	(22)	(22)	81	(40)	(43)
<i>Cervarix</i>	117	(15)	(20)	–	–	–	22	(12)	(8)	95	(16)	(22)
<i>Others</i>	189	26	26	22	(8)	(17)	34	55	45	133	28	32
Vaccines ex COVID	7,873	24	17	4,243	31	18	1,827	27	28	1,803	8	6
Pandemic vaccines	64	(86)	(86)	–	(100)	(100)	57	–	–	7	(97)	(97)
<i>Pandemic adjuvant</i>	64	(86)	(86)	–	(100)	(100)	57	–	–	7	(97)	(97)
Vaccines	7,937	17	11	4,243	22	10	1,884	31	32	1,810	(3)	(5)
HIV	5,749	20	12	3,756	30	17	1,310	10	10	683	–	(3)
<i>Dolutegravir products</i>	5,191	14	6	3,311	19	8	1,239	8	8	641	–	(3)
<i>Tivicay</i>	1,381	–	(7)	823	8	(3)	273	(5)	(4)	285	(14)	(19)
<i>Triumeq</i>	1,799	(4)	(11)	1,217	2	(8)	361	(20)	(19)	221	(8)	(9)
<i>Juluca</i>	636	23	14	494	26	13	127	14	15	15	15	8
<i>Dovato</i>	1,375	75	65	777	82	64	478	58	59	120	>100	>100
<i>Rukobia</i>	82	82	64	79	84	65	3	50	50	–	–	–
<i>Cabenuva</i>	340	>100	>100	294	>100	>100	40	>100	>100	6	>100	>100
<i>Apretude</i>	41	–	–	41	–	–	–	–	–	–	–	–
<i>Others</i>	95	(25)	(29)	31	(37)	(45)	28	(22)	(22)	36	(14)	(17)
Respiratory/Immunology and Other	2,609	29	20	1,830	29	16	366	13	13	413	45	47
<i>Nucala</i>	1,423	25	18	881	28	15	300	17	17	242	24	28
<i>Benlysta</i>	1,146	31	20	949	31	18	83	22	22	114	44	43
<i>Other</i>	40	>100	>100	–	–	–	(17)	–	–	57	>100	>100
Oncology	602	23	17	313	14	3	253	30	31	36	80	75
<i>ZeJula</i>	463	17	12	235	11	–	194	19	20	34	70	75
<i>Blenrep</i>	118	33	25	66	8	(3)	52	86	86	–	–	–
<i>Jemperli</i>	21	>100	>100	13	>100	>100	8	>100	>100	–	–	–
<i>Other</i>	–	–	–	(1)	–	–	(1)	–	–	2	–	–
Specialty Medicines ex COVID	8,960	23	15	5,899	29	16	1,929	13	13	1,132	14	13
Pandemic	2,309	>100	>100	828	38	24	456	>100	>100	1,025	>100	>100
<i>Xevudy</i>	2,309	>100	>100	828	38	24	456	>100	>100	1,025	>100	>100
Specialty Medicines	11,269	37	29	6,727	30	17	2,385	34	35	2,157	69	70
Respiratory	6,548	8	3	3,209	10	(1)	1,384	3	3	1,955	10	9
<i>Arnuita Ellipta</i>	56	19	9	48	20	10	–	–	–	8	14	–
<i>Anoro Ellipta</i>	483	(4)	(9)	233	(16)	(24)	165	11	11	85	10	10
<i>Avamys/Veramyst</i>	321	8	6	–	–	–	65	–	2	256	10	8
<i>Flixotide/Flovent</i>	545	23	15	353	28	16	74	7	7	118	18	16
<i>Incruse Ellipta</i>	196	(4)	(10)	104	(5)	(14)	64	(9)	(7)	28	8	–
<i>Relvar/Breo Ellipta</i>	1,145	2	(2)	498	2	(8)	347	4	4	300	–	2
<i>Seretide/Advair</i>	1,159	(15)	(17)	308	(37)	(43)	287	(11)	(11)	564	3	1
<i>Trelegy Ellipta</i>	1,729	42	32	1,253	47	32	236	18	19	240	47	48
<i>Ventolin</i>	771	7	2	411	5	(5)	116	7	8	244	11	10
<i>Other Respiratory</i>	143	4	6	1	–	–	30	11	7	112	2	5
Other General Medicines	3,570	(1)	(2)	363	10	(1)	695	(14)	(13)	2,512	1	2
<i>Dermatology</i>	376	(6)	(5)	(1)	–	–	107	(18)	(18)	270	–	1
<i>Augmentin</i>	576	35	38	–	–	–	151	22	23	425	41	44
<i>Avodart</i>	330	(1)	(3)	–	–	–	107	(9)	(8)	223	5	–
<i>Lamictal</i>	511	7	1	265	14	3	109	(3)	(3)	137	2	–
<i>Other</i>	1,777	(10)	(10)	99	–	(9)	221	(31)	(31)	1,457	(7)	(6)
General Medicines	10,118	5	1	3,572	10	(1)	2,079	(3)	(3)	4,467	5	5
Total Commercial Operations	29,324	19	13	14,542	22	10	6,348	18	19	8,434	14	14

Financial record continued

Three-year selected financial data

A record of financial performance is provided, analysed in accordance with current reporting practice. The information included in the selected financial data (except for number of employees and adjusted results) is prepared in accordance with International Accounting Standards in conformity with the requirements of the Companies Act 2006 and also with IFRS as issued by the International Accounting Standards Board.

	2023 £m	2022 £m	2021 £m
Group turnover by geographic region			
US	15,820	14,542	11,914
Europe	6,564	6,348	5,370
International	7,944	8,434	7,412
	30,328	29,324	24,696
Group turnover by product group			
Vaccines	9,864	7,937	6,778
Specialty Medicines	10,244	11,269	8,251
General Medicines	10,220	10,118	9,667
	30,328	29,324	24,696
Vaccines turnover			
Shingles	3,446	2,958	1,721
Meningitis	1,260	1,116	961
RSV	1,238	–	–
Influenza	504	714	679
Established Vaccines	3,266	3,085	2,970
Pandemic Vaccines	150	64	447
	9,864	7,937	6,778
Specialty Medicines turnover			
HIV	6,444	5,749	4,777
Respiratory/Immunology and other	3,025	2,609	2,027
Oncology	731	602	489
Pandemic	44	2,309	958
	10,244	11,269	8,251
General Medicines			
Respiratory	6,825	6,548	6,048
Other General Medicines	3,395	3,570	3,619
	10,220	10,118	9,667
Financial results – Total			
Turnover	30,328	29,324	24,696
Profit after taxation from continuing operations	5,308	4,921	3,516
Profit after taxation from discontinued operations and other gains/(losses) from the demerger	–	3,049	1,580
Remeasurement of discontinued operations distributed to shareholders on demerger	–	7,651	–
Profit after taxation from discontinued operations	–	10,700	1,580
Profit after taxation for the year	5,308	15,621	5,096
	pence	pence	pence
Basic earnings per share from continuing operations	121.6p	110.8p	82.9p
Basic earnings per share from discontinued operations	–	260.6p	26.7p
Total basic earnings per share	121.6p	371.4p	109.6p
Diluted earnings per share from continuing operations	119.9p	109.2p	81.8p
Diluted earnings per share from discontinued operations	–	257.0p	26.4p
Total diluted earnings per share	119.9p	366.2p	108.2p

Financial record continued

Three-year selected financial data continued

Financial results – Adjusted	2023 £m	2022 £m	2021 £m
Turnover	30,328	29,324	24,696
Continuing operating profit	8,786	8,151	6,493
Continuing profit before taxation	8,112	7,358	5,774
Continuing profit after taxation	6,855	6,220	4,856

The reconciliation between Total and Adjusted operating profit over the last three years can be summarised as follows:

	2023 £m	2022 £m	2021 £m
Total continuing operating profit	6,745	6,433	4,357
Intangible asset amortisation	719	739	761
Intangible asset impairment	398	296	347
Major restructuring	382	321	424
Transaction-related items	572	1,750	1,143
Divestments, significant legal and other items	(30)	(1,388)	(539)
Adjusted continuing operating profit	8,786	8,151	6,493

The reconciliation between total and Adjusted earnings per share over the last three years can be summarised as follows:

	pence	pence	pence
Total continuing earnings per share	121.6p	110.8p	82.9p
Intangible asset amortisation	13.9p	14.6p	15.2p
Intangible asset impairment	7.5p	5.8p	6.6p
Major restructuring	7.4p	5.9p	8.7p
Transaction-related items	6.9p	34.1p	18.1p
Divestments, significant legal and other items	(2.2)p	(31.5)p	(21.2)p
Adjusted continuing earnings per share	155.1p	139.7p	110.3p
	%	%	%
Return on capital employed	53.0	n/m	25.8

For 2021 and 2023 return on capital employed is calculated as total profit before taxation as a percentage of average net assets over the year and is not restated. Return on capital employed is not calculated for 2022 as it is not meaningful (n/m) as the average net assets over the year included Consumer Healthcare which was demerged on 18 July 2022.

Balance sheet	2023	2022	2021
Non-current assets	40,361	39,377	60,429
Current assets	18,644	20,769	18,674
Total assets	59,005	60,146	79,103
Current liabilities	(21,068)	(22,810)	(23,670)
Non-current liabilities	(25,142)	(27,240)	(34,091)
Total liabilities	(46,210)	(50,050)	(57,761)
Net assets	12,795	10,096	21,342
Shareholders' equity	13,347	10,598	15,055
Non-controlling interests	(552)	(502)	6,287
Total equity	12,795	10,096	21,342

Number of employees	2023	2022	2021
US	12,205	11,946	14,289
Europe	32,675	31,800	38,809
International	25,332	25,654	36,998
	70,212	69,400	90,096
Manufacturing	23,159	23,292	32,141
Selling	26,193	26,310	34,846
Administration	7,888	7,605	11,014
Research and development	12,972	12,193	12,095
	70,212	69,400	90,096

The geographic distribution of employees in the table above is based on the location of GSK's subsidiary companies. The number of employees is the number of permanent employed staff at the end of the financial period. It excludes those employees who are employed and managed by GSK on a contract basis.

Pipelines, products and competition

Pharmaceuticals and Vaccines product development pipeline

Key †	In-license or other alliance relationship with third party	A	Approved
^	ViiV Healthcare, a global specialist HIV company with GSK, Pfizer, Inc. and Shionogi Limited as shareholders, is responsible for developing and delivering HIV medicines	S	Submitted
BLA	Biological Licence Application	Phase I	Evaluation of clinical pharmacology, usually conducted in volunteers
MAA	Marketing Authorisation Application (Europe)	Phase II	Determination of dose and initial evaluation of efficacy, conducted in a small number of patients
NDA	New Drug Application (US)	Phase III	Large comparative study (compound versus placebo and/or established treatment) in patients to establish clinical benefit and safety

MAA and NDA/BLA regulatory review milestones shown in the table below are those that have been achieved. Future filing dates are not included in this list.

Compound	Mechanism of Action/Vaccine Type	Indication	Phase	Achieved regulatory review milestones	
				MAA	NDA/BLA
Oncology					
<i>Ojjaara/Omjjara</i> (mometinib) [†]	JAK1, JAK2 and ACVR1 inhibitor	Myelofibrosis	Approved	A:Jan24	A:Sep23
<i>Jemperli</i> (dostarlimab) [†]	Anti-Programmed Cell Death protein 1 receptor (PD-1) antibody	dMMR/MSI-H 1L endometrial cancer 1L endometrial cancer combination with <i>Zejula</i> (niraparib) Peri-operative dMMR/MSI-H colon cancer Non-small cell lung cancer ¹ Neoadjuvant dMMR/MSI-H rectal cancer	Approved III III II II	A:Dec23	A:Jul23
<i>Zejula</i> (niraparib) [†]	Poly (ADP-ribose) polymerase (PARP) 1/2 inhibitor	1L maintenance ovarian cancer combination with <i>Jemperli</i> (dostarlimab) 1L maintenance non small cell lung cancer (NSCLC) combination with pembrolizumab	III III		
<i>Blenrep</i> (belantamab mafodotin) [†]	ADC targeting B-cell maturation antigen	2L+ multiple myeloma combination with Pomalyst and dexamethasone 2L+ multiple myeloma combination with Velcade and dexamethasone Multiple myeloma in combination with anti-cancer treatments (platform study) 1L multiple myeloma combination with Velcade, Revlimid and dexamethasone	III III II I		
cobolimab [†]	Anti-T-cell immunoglobulin and mucin domain-3 (TIM-3) antibody	Non-small cell lung cancer combination with <i>Jemperli</i> (dostarlimab) and docetaxel	III		
belrestotug [†]	Anti-TIGIT	Non-small cell lung cancer combination with <i>Jemperli</i> (platform study) Squamous cell carcinoma of the head and neck combination with <i>Jemperli</i> and GSK6097608 (platform study)	II II		
4381562 [†]	Anti-PVRIG	Cancer	I		
6097608 [†]	Anti-CD96	Cancer	I		
XMT-2056 ² (wholly owned by Mersana Therapeutics)	STING agonist ADC	Cancer	I		
belantamab	B-cell maturation antigen binder	Multiple myeloma	I		
4524101 [†]	DNA polymerase theta inhibitor	Cancer	I		
5733584 [†]	ADC targeting B7-H4	Gynecologic malignancies	I		
HIV[^]					
<i>Apretude</i> (cabotegravir)	HIV integrase strand transfer inhibitor (long-acting)	HIV pre-exposure prophylaxis HIV infection	Approved I	A:Sep23	A:Dec21
3810109 [†]	HIV broadly neutralising antibody	HIV infection	II		
3739937	HIV maturation inhibitor	HIV infection	II		
4004280	HIV capsid protein inhibitor	HIV infection	II		
4011499	HIV capsid protein inhibitor	HIV infection	II		
4524184 [†]	HIV integrase inhibitor	HIV infection	II		

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Footnotes

(1) non-registrational.

(2) GSK has an exclusive global license option to co-develop and commercialize the candidate.

Pipelines, products and competition continued

Pharmaceuticals and Vaccines product development pipeline continued

Compound	Mechanism of Action/Vaccine Type	Indication	Phase	Achieved regulatory review milestones	
				MAA	NDA/BLA
Infectious Diseases					
Arexvy (RSV vaccine) [†]	Recombinant protein, adjuvanted vaccine	Respiratory syncytial virus prophylaxis in adults 60 years and older	Approved	A:Jun23	A:May23
		Respiratory syncytial virus prophylaxis in older adult population 50-59 years of age	Registration	S:Jan24	S:Dec23
gepotidacin [†]	Triazaacenaphthylene bacterial type II topoisomerase inhibitor	Uncomplicated urinary tract infection (uUTI) Urogenital gonorrhea (GC)	III III		
bepirovirsen [†]	HBV antisense oligonucleotide	Chronic hepatitis B virus infection	III		
<i>Bexsero</i> vaccine	Recombinant protein and outer membrane vesicle vaccine	Prevention of invasive disease caused by <i>N. meningitidis</i> serogroup B in individuals 2 months of age and older (US)	III		
3536819 (Men ABCWY vaccine)	Recombinant protein, outer membrane vesicle, glycoconjugate vaccine	Prevention of invasive disease caused by <i>N. meningitidis</i> serogroups A,B,C,W and Y in adolescents 10-25 years of age	III		
tebipenem pivoxil [†]	Antibacterial carbapenem	Complicated urinary tract infection (cUTI)	III		
ibrexafungerp [†]	Antifungal glucan synthase inhibitor	Invasive candidiasis	III		
ganfeborole [†]	Leucyl t-RNA synthetase inhibitor	Tuberculosis	II		
alpipectir [†]	Ethionamide booster	Tuberculosis	II		
3437949 [†] (Malaria fractional dose)	Recombinant protein, adjuvanted vaccine	Malaria prophylaxis (<i>Plasmodium falciparum</i>)	II		
3536852 [†]	Generalized Modules for Membrane Antigens (GMMA) vaccine	Shigella diarrhea prophylaxis	II		
3528869 [†] (Therapeutic HBV)	Prime-boost with viral vector co- or sequentially administered with adjuvanted recombinant proteins	Treatment of chronic Hepatitis B infections – aims at functional cure by controlling and resolving the clinical sequelae of the infection and reducing the need for further treatment	II		
4023393 (Men ABCWY, 2nd Gen)	Recombinant protein, outer membrane vesicle – conjugated vaccine	Prevention of invasive disease caused by <i>N. meningitidis</i> serogroup A,B,C,W and Y in adolescents and children 6 weeks of age and older	II		
4178116 (Varicella new strain)	Live attenuated vaccine	Active immunization for the prevention of varicella in individuals 12 months of age and older	II		
sanfetrinem cilexetil [†]	Serine beta lactamase inhibitor	Tuberculosis	II		
4106647 [†] (HPV9-AS04)	Recombinant protein-adjuvanted vaccine	Active immunization of girls and women, boys and men (9-45 years), for the prevention of cancer, genital warts and precancerous or dysplastic lesions (girls, boys AIN only) caused by Human papillomavirus (HPV)	II		
4388067 (CHBV ASO combo) [†]	Targeted Immunotherapy (viral vector; adjuvanted recombinant proteins) & Direct Acting Antiviral (GSK's bepirovirsen)	Treatment of chronic Hepatitis B virus infection in individuals >18 years without decompensated cirrhosis	II		
5101955 [†]	MAPS Pneumococcal 24-valent paed	Prevention of pneumonia and invasive pneumococcal disease caused by the <i>Streptococcus pneumoniae</i> 24 serotypes included in the vaccine in children aged 6 weeks – 17 years	II		
5101956 [†]	MAPS Pneumococcal 24-valent	Prevention of pneumonia and invasive pneumococcal disease caused by the <i>Streptococcus pneumoniae</i> 24 serotypes included in the vaccine in adults aged 18 years and older	II		
4406371 (MMRV new strain)	Live attenuated vaccine	Active immunization for the prevention of measles, mumps, rubella, and varicella in children 12 months through 12 years of age	II		

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Pipelines, products and competition continued

Pharmaceuticals and Vaccines product development pipeline continued

Compound	Mechanism of Action/Vaccine Type	Indication	Phase	Achieved regulatory review milestones	
				MAA	NDA/BLA
Infectious Diseases continued					
3993129 (CMV)	Recombinant subunit – adjuvanted vaccine	Cytomegalovirus (CMV) infection prophylaxis in females 16-49 years of age	II		
4382276 [†] (mRNA Seasonal Flu)	mRNA vaccine	Active immunization for the prevention of influenza disease in adults 18 years and older	II		
4396687 [†] (mRNA COVID-19)	mRNA vaccine	Active immunization to prevent COVID-19 disease caused by SARS-CoV-2 in individuals 12 years and older	II		
3943104 [†] (Therapeutic HSV)	Recombinant protein-adjuvanted	Treatment for the suppression of recurrent genital herpes in adults aged 18 years and older	II		
4077164 [†] (iNTS (Typhimurium + Enteritidis))	Bivalent Generalized Modules for Membrane Antigens (GMMA) vaccine	Invasive non-typhoidal salmonella	II		
4077164 [†] (iNTS (S. typhimurium + S. enteritidis + S. Typhi))	Bivalent Generalized Modules for Membrane Antigens (GMMA) vaccine and typhoid conjugate vaccine (TCV)	Invasive non-typhoidal salmonella and typhoid fever	II		
4348413 (Gonorrhoea)	Generalized Modules for Membrane Antigens (GMMA) vaccine	Active immunization for the prevention of gonorrhoea infection in individuals aged 16 to 50 years	II		
daplusiran + tomligisiran	Hepatitis B virus-targeted siRNA sequential combination	Chronic hepatitis B virus infection	II		
3882347 [†]	FimH antagonist	Uncomplicated urinary tract infection (uUTI)	I		
3186899 ^{†3}	CRK-12 inhibitor	Visceral leishmaniasis	I		
3494245 [†]	Proteasome inhibitor	Visceral leishmaniasis	I		
2556286 [†]	Mtb cholesterol dependent inhibitor	Tuberculosis	I		
3923868	PI4K beta inhibitor	Viral COPD exacerbations	I		
3536867 [†] (Salmonella (typhoid + paratyphoid A))	Bivalent Typhoid and Paratyphoid A conjugate	Salmonella typhoid and paratyphoid (A) enteric fever	I		
3965193	PAPD5/PAPD7 inhibitor	Chronic hepatitis B virus infection	I		
5251738 [†]	TLR8 agonist	Chronic hepatitis B virus infection	I		
3772701 [†]	<i>P. falciparum</i> whole cell inhibitor	Malaria	I		
4024484 [†]	<i>P. falciparum</i> whole cell inhibitor	Malaria	I		

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Footnote

(3) Transition activities underway to enable further progression by partner.

Pipelines, products and competition continued

Pharmaceuticals and Vaccines product development pipeline continued

Compound	Mechanism of Action/Vaccine Type	Indication	Phase	Achieved regulatory review milestones	
				MAA	NDA/BLA
Immunology and Respiratory					
<i>Nucala</i> (mepolizumab)	Anti-IL5	COPD	III		
depemokimab [†]	Anti-IL5 (long-acting)	Asthma	III		
		Chronic rhinosinusitis with nasal polyps (CRSwNP)	III		
		Eosinophilic granulomatosis with polyangiitis (EGPA)	III		
		Hypereosinophilic syndrome (HES)	III		
latozinemab [†]	Anti-sortilin monoclonal antibody	Frontotemporal Dementia (FTD) due to Heterozygous Mutations in the Progranulin Gene	III		
		Amyotrophic Lateral Sclerosis (ALS)	II		
		Frontotemporal Dementia (FTD) due to Mutations in the C9orf72 Gene	II		
camlipixant	P2X3 receptor antagonist	Refractory chronic cough	III		
<i>Ventolin</i> , low carbon version of metered dose inhaler	Beta 2 adrenergic receptor agonist	Asthma ⁴	III		
<i>Benlysta</i> (belimumab)	Anti-B lymphocyte stimulator monoclonal antibody	Systemic sclerosis associated interstitial lung disease	II		
3858279 [†]	Anti-CCL17	Osteoarthritis pain	II		
		Diabetic peripheral neuropathic pain	II		
4527226 (AL101) [†]	Anti-sortilin monoclonal antibody	Alzheimer's disease	II		
1070806	Anti-IL18	Atopic dermatitis	II		
3888130 [†]	Anti-IL7	Autoimmune disease	I		
3915393 [†]	Transglutaminase 2 (TG2) inhibitor	Pulmonary fibrosis	I		
5462688 [†]	RNA-editing oligonucleotide	Alpha-1 antitrypsin deficiency	I		
3862995	Anti-IL33	COPD	I		
4347859	Interferon pathway modulator	Systemic lupus erythematosus	I		
Opportunity Driven					
<i>Jesduvroq</i> (daprodustat)	Prolyl hydroxylase inhibitor	Anaemia of chronic kidney disease	Approved		A:Feb23
linerixibat	Ileal bile acid transporter (IBAT) inhibitor	Cholestatic pruritus in PBC (primary biliary cholangitis)	III		
4532990 [†]	HSD17B13 silencer	Non-alcoholic steatohepatitis (NASH)	II		
4172239 [†]	DNMT1 inhibitor	Sickle cell disease	I		

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Footnote

(4) Phase III start expected in 2024.

Pipelines, products and competition continued

Pharmaceutical products, competition and intellectual property

Products	Compounds	Indication(s)	Major competitor brands	Patent expiry dates ¹	
				US	EU
Respiratory					
<i>Anoro Ellipta</i>	umeclidinium bromide/ vilanterol trifenate	COPD	Spiolto/Stiolto RespiMat, Utibron/Ultibro Breezhaler, Duaklir Genuair Bevespi Aerosphere, Brimica Genuair	2027 (NCE) 2027-2031 (device)	2029 (NCE) 2024-2026 (device)
<i>Avamys/Veramyst</i>	fluticasone furoate	Allergic rhinitis	Dymista, Xhance, Nasonex, Fluticasone Gx	expired	expired
<i>Nucala</i>	mepolizumab	Asthma, CRSwNP, EGPA, HES	Fasenra	expired ²	expired ²
<i>Relvar/Breo Ellipta</i>	fluticasone furoate/ vilanterol trifenate	Asthma, COPD	Symbicort, Foster, Budesonide/Formoterol generics, Fluticasone Propionate/Salmeterol generics, Beclomethasone/ Formoterol generics, Ateectura	2025 (NCE) 2027-2031 (device)	2027 (NCE) 2024-2026 (device)
<i>Seretide/Advair</i>	salmeterol xinafoate/ fluticasone propionate	Asthma, COPD	Symbicort, Foster, Budesonide/Formoterol generics, Fluticasone Propionate/Salmeterol generics, Beclomethasone/ Formoterol generics, Ateectura	expired (Diskus device) 2023-2026 (HFA-device)	expired (Diskus device) expired (HFA-device)
<i>Trelegy Ellipta</i>	fluticasone furoate/ vilanterol trifenate umeclidinium bromide	COPD, asthma	Breztri Aerosphere, Trimbow	2027 (NCE) 2027-2031 (device)	2029 (NCE) 2024-2026 (device)
<i>Ventolin</i>	Salbutamol sulphate	Asthma, COPD	Salbutamol/SABA generics, Symbicort as reliever (PRN & MARTI) ³ , Airsupra (US only)	2023-2026 (HFA-device)	expired (HFA-device)
<i>Xevudy</i>	sotrovimab	Early treatment of COVID-19	REGEN-COV, bamlanivimab/ etesevimab, Evusheld	2041	2041
Central nervous system					
<i>Lamictal</i>	lamotrigine	Epilepsy, bipolar disorder	Vimpat, Trokendi XR, Inovelon, <i>Keppra</i> , generics	expired	expired
<i>Keppra</i>	levetiracetam	Epilepsy	Briviact, Vimpat, <i>Lamictal</i> , Depakene, Depacon, generics	NA	NA
Cardiovascular and urogenital					
<i>Avodart & Duodart</i>	dutasteride dutasteride + tamsulosin	Benign prostatic hyperplasia (BPH)	Generics, Finasteride, Alpha Blockers	expired	expired
Anti-bacterials					
<i>Augmentin</i>	Amoxicillin trihydrate/ potassium clavulanate	Common bacterial infections	Generics, Oral Cephalosporins – Cefuroxime axetil, Cefixime, Cefpodoxime, Cefdinir, Cephalexin Oral Macrolides – Azithromycin, Clarithromycin	NA	expired

(1) Unless otherwise stated, patent expiry dates relate to the latest expiring new molecular entity patents in the relevant territory. Where appropriate, these patent expiry dates include granted Patent Term Extensions in the US, granted Supplementary Protection Certificates in multiple countries of the EU and in the UK, and Paediatric Exclusivity periods. Additional exclusivities (for example regulatory data protection or other types of patents) may exist but are not listed in the table.

(2) Regulatory data protection expires 2027 (US) and 2026 (EU).

(3) PRN = use as required *MART = maintenance and reliever therapy.

Pipelines, products and competition continued

Pharmaceutical products, competition and intellectual property continued

Products	Compounds	Indication(s)	Major competitor brands	Patent expiry dates ¹	
				US	EU
Dermatology					
<i>Dermovate, Betnovate, Cutivate, Eumovate</i>	Clobetasol propionate, Betamethasone valerate, Fluticasone propionate, Clobetasone butyrate	Inflammatory skin conditions	Generics, Other topical corticosteroids like Mometasone furoate, Methylprednisolone aceponate and Hydrocortisone	Not marketed in US	expired
Oncology					
<i>Zejula</i>	niraparib	ovarian cancer	Lynparza, Rubraca	2031	2032
<i>Blenrep</i>	belantamab mafodotin	relapsed/refractory multiple myeloma	Abecma, Carvykti, Tecvayli, Talvey, Elrexfio	2032	2032
<i>Jemperli</i>	dostarlimab	dMMR/MSI-H recurrent/advanced endometrial cancer, dMMR solid tumours	Keytruda, Imfinzi+Lynparza	2034	2034
<i>Ojjaara/Omjara</i>	momelotinib	myelofibrosis in patients with anemia	Jakafi, Inrebic, Vonjo	2030	2028
Immuno-inflammation					
<i>Benlysta, Benlysta (SC and IV)</i>	belimumab	systemic lupus erythematosus, lupus nephritis	Lupkynis, Saphnelo	2025	2026
Renal					
<i>Jesduvroq, Duvroq</i>	Daprodustat	anaemia of chronic kidney disease	Evrenzo (roxadustat), vadadustat	2027	Not approved in EU
HIV					
<i>Apretude</i>	Cabotegravir	HIV prevention	Descovy, Truvada	2026	2031
<i>Cabenuva/Vocabria + Rekambys</i>	Cabotegravir, rilpivirine	HIV/AIDS	Descovy, Genvoya, Odefsey, Biktarvy	2026	2031
<i>Rukobia</i>	Fostemsavir	HIV/AIDS	Trogarzo, Sunlenca	2029	2025
<i>Dovato</i>	Dolutegravir, lamivudine	HIV/AIDS	Descovy, Genvoya, Odefsey, Biktarvy	2028	2029
<i>Juluca</i>	Dolutegravir, rilpivirine	HIV/AIDS	Descovy, Genvoya, Odefsey, Biktarvy	2028	2029
<i>Triumeq</i>	Dolutegravir, lamivudine and abacavir	HIV/AIDS	Descovy, Genvoya, Odefsey, Biktarvy	2028	2029
<i>Tivicay</i>	Dolutegravir	HIV/AIDS	Isentress, Prezista Symtuza, Reyataz, Biktarvy	2028	2029

(1) Unless otherwise stated, patent expiry dates relate to the latest expiring new molecular entity patents in the relevant territory. Where appropriate, these patent expiry dates include granted Patent Term Extensions in the US, granted Supplementary Protection Certificates in multiple countries of the EU and in the UK, and Paediatric Exclusivity periods. Additional exclusivities (for example regulatory data protection or other types of patents) may exist but are not listed in the table.

Pipelines, products and competition continued

Vaccine products, competition and intellectual property

Products	Compounds	Indication(s)	Major competitor brands	Patent expiry dates ¹	
				US	EU
<i>Arexvy</i>	Respiratory syncytial virus vaccine	Respiratory syncytial virus vaccination	Abrysvo	2030	2032
<i>Bexsero</i>	meningococcal group-B vaccine	Meningitis group B prophylaxis	Trumenba	2027	2028
<i>Boostrix</i>	diphtheria, tetanus, acellular pertussis	diphtheria, tetanus, acellular Pertussis booster vaccination	Adacel	expired	expired
<i>Infanrix Hexa/ Pediarix</i>	diphtheria, tetanus, pertussis, polio, hepatitis B, Haemophilus influenzae type B (EU)	Prophylaxis against diphtheria, tetanus, pertussis, polio, hepatitis B, Haemophilus influenzae type B (EU)	Pentacel, Pediacel, Pentaxim, Pentavac, Hexaxim, Hexyon Vaxelis	expired	expired
<i>Cervarix</i>	HPV 16 & 18 virus like particles (VLPs), AS04 adjuvant (MPL + aluminium hydroxide)	human papilloma virus type 16 and 18	Gardasil (Silgard)	Not marketed in US	expired
<i>Fluarix Tetra</i>	split inactivated influenza antigens (2 virus subtypes A and 2 subtype B)	seasonal influenza prophylaxis	Intenza, Flumist QIV, Vaxigrip QIV, Fluzone QIV, Fluzone High Dose	expired	expired
<i>FluLaval</i>	split inactivated influenza antigens (2 virus subtypes A and 2 subtype B)	seasonal influenza prophylaxis	Vaxigrip, Mutagrip, Fluzone, Influvac, Aggripal, Fluad, Intenza, Flumist	expired	expired
<i>Menveo</i>	meningococcal group A, C, W-135 and Y conjugate vaccine	Meningitis group A, C, W-135 and Y prophylaxis	Nimenrix, MenQuadfi	2025	2025
<i>Priorix, Priorix Tetraa, Varilrix</i>	live attenuated MMR, Varicella and MMRV vaccines	measles, mumps, rubella and chickenpox prophylaxis	MMR II (M-M-RVaxPro) Proquad, Varivax	expired	expired
<i>Rotarix</i>	Human rotavirus RIX4414 strain	Rotavirus prophylaxis	Rotateq	expired	expired
<i>Synflorix</i>	conjugated pneumococcal polysaccharide	Prophylaxis against invasive disease, pneumonia, acute otitis media	Prevenar (Prevnar)	Note marketed in US	2026
<i>Shingrix</i>	zoster vaccine recombinant, adjuvanted	herpes zoster (shingles)	Zostavax	2029	2031

(1) Unless otherwise stated, patent expiry dates relate to the latest expiring new molecular entity patents in the relevant territory. Where appropriate, these patent expiry dates include granted Patent Term Extensions in the US, granted Supplementary Protection Certificates in multiple countries of the EU and in the UK, and Paediatric Exclusivity periods. Additional exclusivities (for example regulatory data protection or other types of patents) may exist but are not listed in the table.

Principal risks and uncertainties

We outline below the principal risks and uncertainties relevant to GSK's business, financial condition and operations that may affect our performance and ability to achieve our objectives. These are the risks that we believe could cause our actual results to differ materially from expected and historical results.

Operating in the biopharmaceutical sector carries various inherent risks and uncertainties that may affect our business.

We must comply with a broad range of laws and regulations which apply to the research and development, manufacturing, testing, approval, distribution, sales, and marketing of pharmaceutical and vaccine products. These affect the cost of product development, the time required to reach the market and the likelihood of doing so successfully on an uninterrupted basis.

As rules and regulations change, government interpretation and policy evolves, and our business activities develop, the nature of a particular risk may also alter. Changes to regulatory regimes may be substantial. Any alteration in, and failure to comply with, applicable laws and regulations could materially and adversely affect our financial results.

Similarly, our global business exposes us to litigation and government investigations, including product liability litigation, patent and antitrust litigation and sales and marketing litigation.

Litigation and government investigations, and the related provisions we may make for unfavourable outcomes and increases in related costs, such as insurance premiums, could also materially and adversely affect our financial results.

More detail on the status and various uncertainties in our significant unresolved disputes and potential litigation is set out in Note 47 'Legal proceedings' on page 239.

More details regarding our risk management framework and how we identify our principal risks can be found on pages 57 to 61 and incorporated in this section. Other risks related to Environmental, Social, and Governance (ESG) which are not at the level of principal risks, including environmental sustainability and climate change, are managed through our six focus areas, and as described in our ESG Performance Report. Additional information on climate-related risk management is in our climate-related financial disclosure on pages 62 to 70.

UK regulations require a description of principal risks and uncertainties and an explanation of how these are being managed or mitigated. Below is a description of each of our principal risks, together with a summary of their impact and how we manage each risk across our businesses. They are not listed in order of significance and are consistent with the principal risks detailed on pages 59 to 61.

Patient safety

Risk definition

The risk that GSK, including our third parties, fails to appropriately collect, assess, follow up, or report human safety information, including adverse events, from all potential sources or that GSK potentially fails to appropriately act on any relevant findings that may affect the benefit-risk profile of a medicine or vaccine in a timely manner.

Risk impact

GSK will not tolerate an unfavourable benefit-to-risk profile for patients who use our products. As the most important consequence of ineffective pharmacovigilance is the potential for harm to patients, we maintain robust processes for managing human safety information, conducting timely safety signal detection, and ensuring appropriate measures are in place to manage risks to patients. GSK also intends to fully comply with pharmacovigilance and other relevant regulations worldwide. Non-compliance could result in inspection findings, regulatory scrutiny, civil or criminal sanctions and either temporary or permanent loss of product marketing authorisation. We regularly review and respond to all patient safety risks to limit the potential for reputational damage, loss of trust by patients and healthcare providers, product-related litigation, and loss of shareholder confidence.

Context

We are accountable for safeguarding patients and clinical trial participants who receive our medicines and vaccines, whether in development or marketed, from harm. While an unforeseen event that unfavorably shifts the benefit/risk profile is not a probable occurrence, such an event cannot be fully discounted; we mitigate this risk through robust safety evaluation and product risk management activities.

Our Chief Medical Officer is the single point of accountability for benefit/risk decision-making. Cross-functional Safety Review Teams continually assess new safety and efficacy information for every GSK product throughout its life cycle. Our Global Safety Board, under the leadership of our Chief Medical Officer and Head of Global Safety, reviews product safety at established milestones and in every situation where there might be a potential impact on a benefit/risk profile.

We must operate in a complex and restrictive pharmacovigilance regulatory environment, sometimes complicated by variable requirements between regulatory agencies. Such regulatory complexity is further illustrated by instances of regulatory agencies taking decisions on the safety of medicines and vaccines based on externally available data that may not be accessible to the marketing authorisation holder. This trend could inhibit our ability to make timely decisions and take appropriate action in relation to the safety of our products, or to confirm or refute conclusions asserted by external parties. This has the potential to extend beyond regulatory agencies to next-generation digital health data held by technology companies or other data custodians, and inaccessible by our industry and/or regulatory agencies.

Principal risks and uncertainties continued

Patient safety continued

There are many sources of information that might trigger an increase in reporting related to products and/or adverse events (such as media coverage, social media, government health authorities, etc.). Ineffective management of patient safety risks could not only result in reputational damage, loss of trust by patients and healthcare providers, and decline in shareholder confidence, but could also increase the volume of product-related litigation, including class-action lawsuits, which is regularly faced by GSK and our industry in general.

Mitigating actions

Our Chief Medical Officer is accountable for the Patient Safety enterprise risk and human safety matters, in collaboration with the Head of Global Safety. A cross-enterprise safety governance board oversees implementation of our control framework, including risk management. Our Global Safety Board ensures that we address human safety proactively throughout a product's lifecycle. Our global policy on management of human safety information requires that all employees immediately report issues relating to the safety of our products. Our third party risk management framework supports us in identifying and training any third parties who may encounter human safety information.

In 2023, we took additional steps to strengthen how we safeguard patients and enhance the execution of our pharmacovigilance operational activities. We have added risk management and benefit/risk expertise to our Global Safety organisation that will enhance our ability to define the risk management strategy for an evolving portfolio for which more complex risk minimisation measures may be required, to be followed in 2024 with a new system that will better enable us to track the implementation and effectiveness of our risk management plans. We have transitioned to a simplified, more efficient approach for collecting, following up, and reporting human safety information including adverse events from all potential sources. We are continuing to build capability across all GSK staff who hold accountability for our Pharmacovigilance Quality Management System and are in the process of embedding a simplified process for managing pharmacovigilance agreements and safety clauses between GSK and third parties, including our strategic partners.

Product quality

Risk definition

The risk that GSK or our third parties potentially fail to ensure appropriate controls and governance of quality for development and commercial products are in place; compliance with industry practices and regulations in manufacturing and distribution activities; and terms of GSK product licenses and supporting regulatory activities are met.

Risk impact

A failure to ensure product quality could have far-reaching implications for patient safety, cause product launch delays, drug shortages or product recalls, and have regulatory, legal, and financial consequences. These could materially and adversely affect GSK's reputation and financial results.

Context

The external environment for product quality remains challenging. An increase in supplier mergers in our supply network can create challenge in influencing their quality standards. The rapid advancement and use of digital technologies such as artificial intelligence and machine learning (AI/ML) within an evolving regulatory framework introduces both opportunity for modernisation and potential to impact product quality if not adequately controlled. There will be a need to adopt and adapt to new, updated guidance on this as it emerges. The threat of cyber-attacks and data breaches across the industry could risk the integrity of product quality data and its audit trail. Additionally, a gradual divergence in regulatory expectations during inspections, particularly from some health authorities, presents a challenge to our sites as they prepare for inspections. Retaining expertise in biopharma and the deep capability to support digital progression has the potential to be a challenge in a highly competitive environment.

Mitigating actions

We align an extensive global network of quality and compliance professionals from site-level to senior management within each business unit to provide oversight and assist with the delivery of quality performance and operational compliance. We deliver this management oversight through a hierarchy of quality councils and a Global Head of Quality.

We are expanding our Quality Management System, Good Manufacturing Practice Audit and Quality assurance oversight programme across R&D to ensure that we mitigate potential product quality risk across the end-to-end process. We have implemented a risk-based approach to assessing and managing third party suppliers that provide materials used in our finished products including monitoring third party labs and how they are independently checking goods. We expect contract manufacturers that make our products to comply with GSK standards and regularly conduct audits to provide us with assurance. We use key risk indicators to support risk management activities and provide leadership teams and quality councils with an integrated assessment of product quality performance.

Throughout 2023, we continued to actively manage the deployment of plans to align with the New Annex 1 guidance for the manufacture of sterile medicinal products in the context of global equipment and component supply chain constraints effecting the industry. We are increasingly applying advanced digital technologies and insights to drive scientific excellence to enhance and modernise the development, manufacture and testing of our products and to protect our data. We are actively contributing to industry advocacy and to influence thinking on the regulatory frameworks for these advancing technologies to support patient safety benefit and access.

Principal risks and uncertainties continued

Product quality continued

We are collaboratively working with other pharma companies and industry trade associations to respond to questions from the EMA and completing the safety evaluation of the use of Titanium Dioxide in medicines, as well as identifying potential substitutes. We are working with industry to monitor emerging risk factors and regulatory intelligence and guidance on Nitrosamines.

We continue to adapt our procedures to the evolving expectations on this topic and work on our mitigation plans alongside regular engagement with regulators, at all times ensuring our inspection readiness for all the markets we serve.

Financial controls and reporting

Risk definition

The risk that GSK fails to comply with current tax laws; fails to report accurate financial information in compliance with accounting standards and applicable legislation; or incurs significant losses due to treasury activities.

Risk impact

Non-compliance with existing or new financial or new ESG reporting and disclosure requirements, or changes to the recognition of income and expenses, could expose GSK to litigation and regulatory action and could materially and adversely affect our financial results. Failure to comply with changes in the substance or application of the laws governing transfer pricing, dividends, tax credits and intellectual property could also materially and adversely affect our financial results. Failure to comply with applicable laws and regulations could result in GSK being investigated by relevant government agencies and authorities and/or in legal proceedings against us. Government investigations and litigation, can be unpredictable and regardless of their outcome, may be costly, require significant management attention, and damage our reputation. Inconsistent application of treasury policies, transactional or settlement errors, or counterparty defaults could lead to significant losses.

Context

We are required by the laws of various jurisdictions to publicly disclose our financial results and any events that could materially affect the Group's financial results. Regulators routinely review the financial statements of listed companies for compliance with new, revised, or existing accounting and regulatory requirements. We believe that we comply with the appropriate regulatory requirements concerning our financial statements and the 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 could lead to restatements of previously reported results and significant penalties. Our Treasury group deals daily in high value transactions, mostly foreign exchange, and cash management transactions. These transactions involve market volatility and counterparty risk. The Group's effective tax rate reflects the locations of our activities and the value they generate, which determine the jurisdictions in which profits arise and the applicable tax rates.

These may be higher or lower than the UK statutory rate and may reflect regimes that encourage innovation and investment in R&D by providing tax incentives which, if changed, could affect GSK's tax rate. In addition, the worldwide nature of our operations means that our cross-border supply routes, necessary to ensure supplies of medicines and vaccines, can result in conflicting claims from tax authorities as to the profits to be taxed in individual countries.

This can lead to double taxation, with profits taxed in more than one country. The complexity of tax regulations also means that we may occasionally disagree with tax authorities on the technical interpretation of a particular area of tax law. The tax charge included in our financial statements is our best estimate of tax liability pending any audits by tax authorities. We expect there to be a continued focus on tax reform, driven by initiatives by the OECD and the EC to address the tax challenges arising from digitalisation of the economy.

Together with domestic initiatives around the world, these may result in significant changes to established tax principles and an increase in tax authority disputes. Regardless of their merit or outcomes, these may be costly, divert management attention and adversely impact our reputation and relationship with key stakeholders. Laws, regulations, orders and other measures restrict dealings with certain countries, governments, government officials, entities, individuals, and the use of financial institutions and movement of funds.

Mitigating actions

We keep up to date with the latest developments in financial reporting requirements by reviewing updates from regulators, working with our external auditor and legal advisors and performing and responding to emerging risks. Financial results are reviewed and approved by regional management, before being reviewed by GSK's Group Financial Controller and Chief Financial Officer (CFO). This allows our Financial Controller and CFO to assess the evolution of the business over time, and to evaluate its performance to plan. Significant judgements are reviewed and confirmed by senior management. We integrate technical or organisational transformation, newly acquired activities and external risks into our risk assessments and apply appropriate controls and reviews.

We maintain a control environment designed to identify material errors in financial reporting and disclosure. We have a standardised global financial reporting operating model. The design and operating effectiveness of key financial reporting controls are regularly reviewed by management and tested by external third parties. The few locations which are not on the standard model apply a minimum standard set of controls which are reviewed by management and monitored independently. This gives us assurance that controls over key financial reporting and disclosure processes are operating effectively. Our Global Finance Risk Management and Controls (FRMC) group provides extra support during significant transformations, such as system deployment or management/structural reorganisations. We add operational resources and adapt programme timelines to ensure processes and controls are maintained during significant changes. The Disclosure Committee, reporting to the Board, reviews GSK's quarterly results and annual report.

Principal risks and uncertainties continued

Financial controls and reporting continued

Throughout the year, in consultation with its legal advisors, the Disclosure Committee also determines whether it is necessary to disclose publicly information about the Group through stock exchange announcements. The Treasury Management Group meets regularly to ensure that liquidity, interest rate, counterparty, foreign currency transaction and foreign currency translation risks are all managed in line with the prudent approach detailed in the risk strategies and policies adopted by our Board. Counterparty exposure is subject to defined limits approved by the Board for both credit rating and individual counterparties.

The Middle Office within Treasury monitor the management of counterparty risk in line with agreed policy with oversight from a corporate compliance officer, operating independently of Treasury. Further details on mitigation of Treasury risks can be found on pages 243 to 245. We manage tax risk through robust internal policies, processes, training, and compliance programmes. We maintain open and constructive relationships with tax authorities worldwide. We monitor government debate on tax policy in our key jurisdictions, so that we can understand any potential future changes in tax law and share an informed point of view.

Where relevant, we provide pragmatic and constructive business input to tax policy makers, either directly or through industry trade bodies. This includes advocating reform to support economic growth and job creation, as well as the needs of our patients and other key stakeholders. Our tax affairs are managed on a global basis by a team of tax professionals, led by the Global Head of Tax, who work closely with the business on a day-to-day basis. The Global Tax team is suitably qualified for the roles they perform, and we support their training needs so they can provide up to date technical advice in line with their responsibilities. We submit tax returns according to statutory time limits and engage proactively with tax authorities to ensure our tax affairs are current, entering into continuous audit programmes and advance pricing agreements where appropriate. These arrangements provide long-term certainty for both tax authorities and GSK over the tax treatment of our business, based on full disclosure of all relevant facts. We seek to resolve any differences of interpretation in tax legislation with tax authorities in a cooperative manner. In exceptional cases, we may have to resolve disputes through formal proceedings.

Legal matters

Risk definition

The risk that GSK or our third parties potentially fail to comply with certain legal requirements for the development, supply and commercialisation of our products and operation of business, and specifically in relation to requirements for competition law, anti-bribery and corruption, and sanctions. Any failure to meet compliance and legal standards for these particular areas could lead to increasing scrutiny and enforcement from government agencies.

Risk impact

Failure to mitigate legal risk could expose GSK and associated persons to governmental investigation, regulatory action, and civil and criminal liability. It may compromise GSK's ability to supply its products under certain government contracts. In addition, failure to manage legal risk could have substantial implications for GSK's reputation and the credibility of senior leaders. It might erode investor confidence in our governance, risk management and future performance, and have a consequential negative impact on share performance. It could also lead to the imposition of significant financial penalties and the imposition of additional reporting obligations.

Context

The overall environment for anti-bribery and corruption, competition law and sanctions and export controls remains challenging. There continues to be a strong enforcement appetite for bribery investigations and prosecutions, with a particular focus on the conduct of multinational companies wherever they operate. The focus on sanctions, export controls and competition law enforcements has increased. From a sanctions perspective, we have seen penalties for violations levied on companies from a number of different industries. Merger control has seen increasing intervention with greater divergence in decisions and policy by enforcement agencies. Financial penalties handed down in these types of case are often very significant.

Supportive aspects of the external environment include an increase in focus on corporate transparency. Advances in technology and the use of data analytics are also providing better platforms to streamline processes and detect potential issues.

Mitigation actions

Our Group General Counsel oversees and is accountable for the Legal Matters principal risk. We have enterprise anti-bribery and corruption, competition law and sanctions control frameworks and programmes designed to ensure compliance with applicable laws and regulations, building on our Code, culture and business standards, and monitor and adapt to evolving regulations and our business activities. Our programmes include senior leader commitment, setting the tone at the top.

These control frameworks are based on globally recognised and accepted principles and include global policies, written standards and controls to govern business activities that give rise to these risks. We mandate enhanced controls, including due diligence requirements and sanctions screening, for specific high-risk activities such as interactions with government officials, during business development transactions and engagement with third parties.

We regularly provide anti-bribery and corruption, competition law and sanctions training to employees, and relevant complementary workers and third parties in accordance with their roles, responsibilities and risks they face. We include aspects of these key risks in our annual mandatory training and reinforce to our workforce clear expectations regarding acceptable behaviours.

We leverage data analytics and use information from our monitoring and other assurance activities, key risk indicators, investigations, and Speak Up channels to identify specific areas for intervention, and drive continuous improvements and enhancements to our controls. We investigate allegations of non-compliance and take disciplinary action as required and where permitted locally.

Principal risks and uncertainties continued

Legal matters continued

Dedicated teams are responsible for the implementation and evolution of the risk framework and programmes for anti-bribery and corruption, competition law, and sanctions.

We continuously assess, monitor and understand our risk exposure to related risks, including our money laundering risk, and actively consider and implement improvements to the risk framework and programmes based on internal and external learnings, considering the complexity and geographic breadth of the risk.

Commercial practices

Risk definition

The risk that GSK or our third parties potentially engage in commercial activities that fail to comply with laws, regulations, industry codes, and internal controls and requirements.

Risk impact

Failure to engage in activities that are consistent with the letter and spirit of the law, industry regulations, or the Group's requirements relating to sales and promotion of medicines and vaccines; with appropriate interactions with healthcare professionals (HCPs), organisations and patients; with legitimate and transparent transfers of value; and with pricing and competition (or antitrust) regulations in commercial practices, including trade channel activities and business tendering, could materially and adversely affect our ability to deliver our strategy and long-term priorities. Additionally, it may result in incomplete awareness of the risk/benefit profile of our products and possibly suboptimal treatment of patients and consumers; 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. Any practices that are found to be misaligned with our culture could also result in reputational harm and dilute the trust established with external stakeholders.

Context

We operate in a highly regulated and extremely competitive biopharma industry, amongst peers who make significant product innovations and technical advances and intensify price competition. Additional external factors include access limitations to our customers, macroeconomic inflationary dynamics, and pricing pressure across markets. To achieve our strategic objectives, we must continue to develop commercially viable new products, sustain reliable supply, and deliver additional uses for existing products that address the needs of patients, consumers, HCPs and payers.

Financially, new products/indications carry with them an uncertainty of future success. Product development is costly, lengthy, and uncertain, and carries the potential for failure at any stage. Even after successful product development, we face challenges in how we launch, and our competitors' products or pricing strategies could render our assets less competitive. We support product innovation through our continued focus on both in-person and virtual engagement, with a constant focus on our patient. Once we have an approved medicine or vaccine, it is our obligation to provide important information to the healthcare community in various ways, always in a responsible, legal, and ethical manner.

Appropriate product promotion ensures HCPs have access to the information they need, that patients and consumers have the facts about the medicines and vaccines they require, and that products are prescribed, recommended, or used in a manner that provides healthcare benefit. We are committed to the ethical and responsible commercialisation of our products in support of our purpose to improve the quality of human life and get ahead of disease together.

Mitigating actions

We have evolved policies and standards incrementally to ensure that commercial activities that we undertake or are conducted on our behalf are executed within our established governance. We train employees on relevant information with a focus on interactive learning and elements of behavioural science. All our commercial activities worldwide must conform to high ethical, regulatory, and industry standards. Where local standards differ from global ones, we apply those that are most stringent. Where the standards of an acquired company or joint venture partner differ from our global standards, we remediate legacy policies and implement revisions, so they align. Our businesses continue to use our internal control framework to support the assessment and management of risks.

Principal risks and uncertainties continued

Commercial practices continued

Business unit risk management and compliance boards, which manage risks across global and in-country business activities, oversee commercial activities and their monitoring programmes. All promotional materials and activities must be reviewed and approved according to our policies and standards and conducted in accordance with local laws and regulations; these requirements seek to ensure that such materials and activities fairly represent the Group's products or services. Where necessary, in the event of misconduct, we have disciplined employees, up to and including termination of contract, and, applied/enforced GSK's senior leader recoupment policy. We have continued to evolve our incentive programme for sales representatives to better recognise and reward individual effort. In nearly all markets, the capped variable pay element of representatives' compensation is evaluated on the basis of individual sales targets.

We allow fair-market value payments to be made by GSK to expert practitioners to speak about our innovative medicines and vaccines during a restricted period in a product's lifecycle, or when new and competitive data is published. To support this, in 2023 we embedded a global end-to-end expert engagement process, rolled out a Healthcare Organisation (HCO) process, created a new standard operating procedure for tenders, updated our External Expert Engagement operating procedure, and further strengthened our interactive digital media channel controls through the identification of all channels, and the contracting of a third party to monitor these channels across GSK, to drive consistent ways of working and efficiencies and strengthen controls through automation and use of data. Where permitted we report payments to individual HCPs as part of our commitment to transparency and responsible disclosure.

Scientific and patient engagement

Risk definition

The risk that GSK or our third parties potentially fail to engage externally to gain insights, educate and communicate on the science of our medicines and associated disease areas, and provide grants and donations in a legitimate and transparent manner compliant with laws, regulations, industry codes and internal controls and requirements.

Risk impact

Without controls in place, the risk could result in real, perceived, or disguised promotion including off-label and prior authorisation promotion, and real or perceived provision of medical advice. This could lead to reputational damage, competitor complaints, regulatory inspections with subsequent corrective actions, or civil litigation.

We must fully and appropriately engage externally to bring patient benefit, and to advance science and innovation, while delivering our strategy. Otherwise, we risk reducing the trust of the public, patients, healthcare professionals, payers, regulators, and governments.

Context

Scientific and patient engagements are diverse non-promotional activities directed at healthcare professionals, patients, payers, and external stakeholders. Such engagements aim to improve patient care through the exchange or provision of knowledge on the use of our products and related diseases. Scientific and patient engagement with external stakeholder groups is vital to GSK, as a research-based biopharma company that is ambitious for patients and to advance science and medicine.

We expect our activities to be scientifically sound and accurate, conducted ethically and transparently, and compliant with applicable codes, laws, and regulations. There are many industry and local codes and laws and other regulations that apply (such as Privacy or Data integrity). That means measured risk-taking, rooted in sound ethical considerations, and principles-based decision-making, training, communication, and monitoring of such activities are key to managing the risk and enabling full and appropriate engagement.

Mitigating actions

Our Chief Medical Officer (CMO) oversees all non-promotional scientific and patient engagement (SPE) as enterprise risk owner. The GSK Code of Practice is the key internal policy for non-promotional engagement activities. These activities include scientific interactions, support for medical education, advice seeking, gathering insights on unmet needs of patients, scientific communication of our research, and disease awareness, healthcare support services and patient support programs.

Process simplification continued into 2023. Global process owners accountable for the end-to-end process have been assigned for the simplified processes: Seeking Advice, Content Approval, Medical Information and Medical Education. This accountability includes the comprehensive oversight of the process, the creation of an appropriate internal control framework and continuous evaluation of process for improvement where necessary.

All SPE materials and activities must be reviewed and approved according to our policies and standards. Additional controls for the review of SPE content have been implemented to continue to ensure content is non-promotional, accurate, fair, objective and balanced and will not be perceived as promotion.

We have further modernised our digital approach to HCPs, embedded our framework for interactions with patients and patient organisations, and developed our policy for healthcare support services and patient support programmes and applied our internal principles to these activities. An internal framework for Software as Medical Device was established including an Expert Panel that provides business owners with multi-disciplinary advice. The cross-business unit SPE risk council oversees SPE activities and reviews monitoring and audit data, while the SPE network reviews the maturity of the internal control framework of the SPE processes. We continuously improve our internal controls, systems and networks to identify emerging risks early and to support staff to conduct activities in compliance with GSK's culture and policies and local laws and regulations while building effective risk management and management monitoring systems.

Principal risks and uncertainties continued

Data ethics and privacy

Risk definition

The risk that GSK or our third parties potentially fail to ethically collect; use; re-use through artificial intelligence, data analytics or automation; secure; share and destroy personal information in accordance with laws, regulations, and internal controls and requirements.

Risk impact

Non-compliance with data privacy laws globally could lead to harm to individuals and GSK. It could also damage trust between GSK and individuals, communities, business partners and government authorities. Many countries have increased the enforcement powers of their data protection authorities by allowing them to impose significant fines, restrict cross-border data flows, or temporarily ban data processing. Many new national laws also enable individuals to bring collective legal actions against companies such as GSK for failing to follow data privacy laws.

Context

Data protection and privacy legislation is diverse, with limited global harmonisation or simplification, making it challenging for multinationals to standardise their approach to compliance. Governments are enforcing compliance with data protection and privacy laws more rigorously.

The approach and focus of data protection and privacy regulators also differs between regions and countries, which creates further challenges for global organisations seeking to implement a single harmonised global privacy programme.

Increases in the volume of data processed and advances in technology have resulted in a greater focus on data governance and the ethical use of personal information, over and above compliance with data privacy laws. Companies seeking to foster innovation in artificial intelligence and other new technologies are faced with evolving decisions from global policymakers on how best to promote trust in these systems and avoid unintended outcomes or harmful impacts.

Additionally, there are a number of emerging laws concerning the localisation of data, restrictions on international transfers and data security, which are changing existing frameworks that GSK has previously relied upon. This increasing trend for data sovereignty affects our ability to drive medical innovation and to effectively operate internationally.

Global regulators (such as the EU, UK, US and China) are also in the process of introducing legislation around the use of artificial intelligence and machine learning (AI/ML). There continues to be considerable uncertainty around the final version of these proposed laws.

Mitigating actions

Our General Counsel is GSK's Enterprise Risk Owner (ERO), and chairs our Digital and Privacy Governance Board, which oversees GSK's overall data ethics and privacy operating model. Each GSK business area has appointed a risk owner accountable for overseeing its privacy risks, supported by privacy leaders within their business. In countries where local data privacy laws require the appointment of a Data Protection Officer (DPO), GSK has made such appointments, including an EU DPO. As a result of GSK's focus on technology, data-driven science, use of AI/ML and our evolving global data strategy, the ERO has appointed a Head of Digital, Privacy and Cybersecurity (Head of DPC), who has day-to-day accountability for designing and implementing the control framework.

The Head of DPC leads a global, cross-functional core team of digital- and privacy-qualified attorneys and compliance professionals, supported by a network of privacy leaders within business units/functions, privacy contacts locally, and the wider Legal & Compliance team. GSK has a global data ethics and privacy framework based on the EU General Data Protection Regulation, which is deployed in every market based on factors including the robustness of local privacy legislation, established data protection authorities, and GSK's footprint.

Beyond those countries, we have deployed a proportionate control framework to set up minimum privacy standards irrespective of any applicable legislation.

Our core team is responsible for:

- operating and improving the centralised global data ethics and privacy control framework
- continuously assessing and providing relevant and proportionate controls and aid to non-deployed markets
- monitoring new, or changing, laws and adapting the privacy framework accordingly
- deploying a comprehensive training programme to drive greater awareness and accountability for managing personal information across the entire organisation
- legal and regulatory expertise in emerging technologies, including artificial intelligence and machine learning

We ensure key GSK privacy network roles have sufficient training and experience to carry out their roles effectively. We continuously improve our processes, such as issue identification, reporting and handling, through monitoring. Our core team works with the business to ensure we build in privacy controls into all existing and new business initiatives, as well as ensuring we meet our accountability obligations in accordance with global data protection and privacy laws.

Principal risks and uncertainties continued

Research practices

Risk definition

The risk that GSK or our third parties potentially fail to adequately conduct ethical and credible pre-clinical and clinical research, collaborate in research activities compliant with laws, regulations, and internal controls and requirements.

Risk impact

The potential 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 GSK by governmental and private plaintiffs (product liability suits and claims for damages), loss of revenue due to inadequate patent protection or inability to supply our products, and regulatory action such as fines, penalties, or loss of product authorisation. Poor data integrity and governance could compromise GSK's R&D efforts and negatively impact our reputation. Any of these could materially and adversely affect our financial results and damage the trust of patients and customers.

Context

Research involving animals can raise ethical concerns. In many cases, however, research involving animals is the only way to investigate the effects of a potential new medicine in a living body other than in humans. Animal research provides critical information about the causes and mechanisms of diseases and therefore remains a vital part of our research. We continually seek ways in which we can minimise our use of animals in research, development, and testing, while complying with regulatory requirements and reducing the impact on the animals used. Human subject research is critical to assessing and demonstrating the safety and efficacy of our investigational products or further evaluating our products once they have been approved.

This research includes clinical trials in healthy volunteers and patients and adheres to regulations and high ethical, medical, and scientific standards. We disclose the results of this research externally regardless of whether they reflect positively or negatively on our products, so that the scientific community can learn from the outcomes of our research. We also work with human biological samples which are fundamental to the discovery, development, and safety monitoring of our products.

We are committed to managing human biological samples in accordance with relevant laws, regulations, and ethical principles, and in a manner that respects the interests of sample donors. Data is pivotal to our R&D strategy, and we are maximising the use of data to serve patients. Governing our data in accordance with relevant laws, regulations, contractual obligations, expectations, and our culture across data ethics, privacy, information and cyber security, and data integrity is essential.

We use a wide variety of biological materials in the discovery, research, and development of our assets. We are committed to ensuring research is compliant with terms and conditions of licenses, agreements or authorisations under which we acquire, use, or transfer biological materials and technologies. 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 research and development. We support the equitable access and fairness principles of access and benefit sharing (ABS) outlined in the CBD and the Nagoya Protocol. We also recognise the importance of appropriate, effective, and proportionate implementation measures at national and regional levels.

Mitigating actions

The Research Practices risk is overseen by an enterprise framework that seeks to strengthen governance across R&D. Under the leadership of the Research Practices Enterprise Risk Owner, management of the risk takes a pragmatic approach to information sharing, streamlining risk identification and escalation while ensuring ownership of risk mitigation stays with the business.

We have an established Office of Animal Welfare, Ethics and Strategy and Risk (OAWESR), led by our Chief Veterinary Officer, which oversees and ensures the humane and responsible care and use of animals, the conduct of ethical reviews and independent scientific reviews of animal studies, and advocates for the application of non-animal alternatives. The OAWESR provides a framework of animal welfare governance; defines and provides oversight for animal care and use programmes; promotes the replacement, refinement, and reduction of animal use in research; conducts quality assessments and manages a programme of due diligence of external animal research.

Ensuring we implement and maintain proper data governance controls remains an important priority, especially as our scientific strategy is evolving to take advantage of the breadth of our data (for example: genomics and artificial intelligence and machine learning (AI/ML)). We focus on building data integrity, privacy and usage controls into our internal control framework. Quality assurance teams conduct audits to provide independent business monitoring of our internal controls. Our R&D organisation maintains and controls pre-publication procedures to guard against public disclosure before patent applications are filed. In addition, because a lack of data integrity in preparing patent application data and information can lead to a loss of patent protection, legal experts collaborate with R&D to support the review process for new patent applications. Our R&D organisation also collaborates with legal experts throughout the development of our assets to take account of any relevant third-party patent rights.

Principal risks and uncertainties continued

Environment, health, and safety (EHS)

Risk definition

The risk that GSK or our third parties potentially fail to ensure appropriate controls and governance of the organisation's assets, facilities, infrastructure, and business activities, including execution of hazardous activities, handling of hazardous materials, or release of substances harmful to the environment that disrupts supply or harms employees, third parties or the environment.

Risk impact

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

Context

GSK is subject to the health, safety and environmental laws of various jurisdictions. These laws impose duties to protect people, the environment and the communities in which we operate.

Mitigating actions

The GSK Leadership Team is responsible for EHS governance and risk oversight. They ensure there is an effective control framework 'in-place' and 'in-use' to manage the EHS risks, impacts, and legal compliance issues in each of our businesses. This includes assigning responsibility to senior managers for providing and maintaining our controls and for ensuring that tiered monitoring and governance processes are in place within their business units.

Function leaders ensure that the EHS control framework is implemented effectively in their respective business area, that it is compliant with applicable laws and regulations, and that it is adequately resourced, maintained, communicated, and monitored. Every employee and qualified contractor acting on behalf of GSK is personally responsible for ensuring that they follow all applicable local standard operating procedures. Our risk-based, proactive approach is articulated in our global EHS policy and detailed in our global EHS standards, against which we audit all our operations to ensure compliance. We ensure hazards are appropriately controlled through the design of facilities, equipment and systems. These rigorous procedures, when applied correctly, put effective barriers in place to protect employees' health and safety.

We have refreshed and rebranded the 12 Life Saving Rules across GSK, with global initiatives to embed the rules in daily operations. Our Safety Leadership Experience training continues across GSK, using incident knowledge to enhance learnings and build a strong, collaborative safety culture. Our Contractor Safety assessment is being deployed with the support of an external expert review of current GSK contractor management. We are improving driver safety through safer cars and enhanced training.

Information and cyber security

Risk definition

The risk that GSK or our third parties potentially fail to ensure appropriate controls and governance to identify, protect, detect, respond, and recover from cyber incidents through unauthorised access, disclosure, theft, unavailability or corruption of GSK's information, key systems, or technology infrastructure in accordance with applicable laws, regulations, industry standards, internal controls and requirements.

Risk impact

Failure to adequately protect our information and systems against cyber security threats may cause harm to patients, workforce and customers, disruption to our business and/or loss of commercial or strategic advantage, regulatory sanction, or damage to our reputation.

Context

The external environment continues to be extremely challenging, making it hard to keep pace with increasingly sophisticated cyber security threats. Factors include increased geopolitical conflict and digital nationalism, rising frequency and severity of data breaches and the growing capability and sophistication of cyber threat actors with additional tools like generative AI to propagate their attacks. GSK's business relies on operating a highly connected information network of internal and external systems which hold confidential research and development, manufacturing, commercial, workforce and financial data. This means that our systems and information have been and will continue to be targeted by cyber security threat actors. Acceleration in the use of digital, data and analytics, AI/ML and cloud computing capabilities to drive GSK's pipeline, performance and productivity requires us to continuously adapt and strengthen our controls and defensive capabilities.

Principal risks and uncertainties continued

Information and cyber security continued

GSK also relies on third-party contractors, partners and suppliers who face similar cyber security threats, which emphasises our focus on third party risk management. Additionally, hybrid working environments create a larger and more complex attack surface for cyber security threat actors to exploit. With employees accessing company resources from various locations and devices, new threats and vulnerabilities could arise.

Mitigating actions

How we manage cyber security risk

We use our corporate enterprise risk management and internal control framework to manage and oversee our Information and Cyber Security principal risk, and we follow our corporate governance hierarchy for risk reporting and escalation. Our Chief Information Security Officer (CISO) heads our Cyber Security Office and is responsible for identifying and putting in place controls and measures to help GSK mitigate and manage cyber security risks. This includes actively monitoring and initiating remediation or other actions to respond to cyber security intelligence and threats. It also includes ongoing investment in people, process and technology to improve our ability to prevent, detect, respond to and recover from any cyber security incidents. We monitor this risk using key risk indicators which include tolerance thresholds reported monthly to the business and quarterly through the governance channels.

We also have a third-party security risk management programme to assess cyber security risk when selecting and onboarding third parties like external partners and suppliers. We use widely accepted standards and frameworks to benchmark our internal environment and controls and help define our security objectives and desired security outcomes. While our standards and frameworks can evolve in response to our dynamic threat environment, we also rely on external frameworks including:

- the National Institute of Standards and Technology (NIST) Cyber Security Framework for measuring the overall cyber readiness and maturity
- the International Organisation for Standardisations (ISO) 27001/27002 for general information technology controls
- Sarbanes-Oxley (SOX) for assessment of internal controls

We also draw on third-party consultants' expertise in processes for assessing, identifying and managing cyber security risks. This year, our cyber security maturity programme, designed to reduce the risk of our data being compromised, has improved our security posture and our ability to detect, protect against, respond to and recover from malicious cyber activity. We also created an AI Governance Council, which includes the CISO, to assess and manage information security risks around adopting and scaling up AI at GSK.

Information and Cyber Security Governance

The Chief Digital and Technology Officer (CDTO) leads the Digital and Technology function, which includes the CISO and Cyber Security Office. The CDTO is the enterprise risk owner for our Information and Cyber Security principal risk, responsible for managing and reporting on the risk, and the enterprise risk plan. This plan includes a description of the risk, its context, our assessment and risk appetite, how we treat the risk and what actions we need to take to manage it in line with our corporate internal control framework. The CISO is responsible for risk coordination across the organisation, developing and overseeing the implementation of controls, and monitoring and reporting on the enterprise risk plan. Both the Board and the Audit & Risk Committee oversee our cyber security risk. The Risk Oversight and Compliance Council helps the Audit & Risk Committee to oversee the cyber security risks, and our strategies to address them. The CISO reports on cyber security risks throughout the year to the CDTO, Risk Oversight and Compliance Council and the Audit & Risk Committee. This reporting covers, external insights, key risk indicators, management actions, updates on implementing the enterprise risk plan, progress on the cyber maturity programme, and escalations. The Cyber Security Office analyses potential cyber security incidents, supported by internal experts, and gives updates to the CISO. The CISO escalates any cyber incidents with potential for material impact to the Chief Compliance Officer and the CDTO, who in turn escalates to the GSK Leadership Team and Company Secretary, triggering review by the Disclosure Committee to determine materiality. Any material cyber security incidents are subsequently escalated to the Board and Audit & Risk Committee.

Cyber Security Awareness, Training and Readiness

Our cyber security awareness and training programmes include phishing simulations, monthly awareness campaigns and mandatory annual refreshers for all employees, new hires and high-risk roles. We run quarterly phishing simulation tests and related remedial trainings. We also offer optional training and an annual global event. These efforts aim to increase cyber security awareness and foster a culture that security is everyone's responsibility. Also, we run periodic crisis simulation exercises for targeted functions to test our response to cyber security incidents.

Compliance with various governmental cyber security regulations

Our Cyber Security Office, guided by our General Counsel, works to stay abreast of emerging government regulations, trends, and compliance expectations regarding cyber security. As new regulatory guidance becomes available (including the U.S. Securities and Exchange Commission's rules on cyber security related disclosures), we respond with remedial compliance-related actions.

Principal risks and uncertainties continued

Supply continuity

Risk definition

The risk that GSK or our third parties potentially fail to deliver a continuous supply of compliant finished product or respond effectively to a crisis incident in a timely manner to recover and sustain critical supply operations.

Risk impact

.We recognise how important the continuity of supply of our products is to the patients who rely on them. Supply disruption can lead to:

- Product shortages and product recalls
- Regulatory intervention
- Reputational harm
- Lost sales revenue

Consequently, we need sophisticated end-to-end supply chain management with robust crisis management and business continuity plans in place to respond.

Context

We operate our supply chains in a continually evolving, highly regulated environment. There is no single set of global regulations which governs the manufacture and distribution of medicines, and we must adhere to the requirements in all those markets in which we licence, sell or manufacture our products. We rely upon our internal Quality Management System and our Internal Control Framework to ensure we maintain our licence to operate. Our complex end-to-end supply chains often involve third party suppliers, from Active Pharmaceutical Ingredient (API) manufacturers and raw material suppliers through to Third Party Logistics Providers and contract engineering firms. We have integrated risk management into our sourcing and day to day business processes, with emphasis on our Third-Party oversight. External factors continued to challenge supply continuity in 2023. The difficulties with sourcing bioscience materials has eased through the year.

There is a new constraint with third party sterile manufacturing capacity which increases global competition for contract manufacturing operations. We continue to operate our global supply chains in a rapidly changing geopolitical environment. Increasing nationalism and friction between the US and China creates divergence from global supply strategy. We have reacted to this by designing supply routes that de-risk sourcing decisions and use business continuity planning to mitigate and maintain supply continuity, e.g. dual sourcing for materials and adapting supply routes to meet regulatory expectations for both the commercial and late stage clinical supply chains.

Our supply chain imperatives focus on accelerating innovation with the use of technology and data to transform the way we manufacture and supply our medicines and vaccines. We drive our competitive advantage through our long-term strategic partnership with R&D. We focus our talent on the skills needed for the future, addressing skills in new technologies and modalities. We have brought the Vaccines and Medicines supply chains together into one Global Supply Chain organisation to leverage the benefits of our highly skilled workforce. Continual business monitoring is in place to assess the sector-wide risk of the spread of industrial relations challenges arising from global cost of living pressures. Keeping our patients supplied with their medicines is our priority.

Mitigating activities

Risk management

Our Medicines and Vaccine supply chains are set up to ensure sustainable global supply. The GSK Internal Control Framework drives our approach to risk management, and it has been designed to identify emerging new risks and support clear decision making. Risk oversight is managed through a hierarchy of Risk Management and Compliance Boards to assure risk mitigation (including identifying new and emerging threats).

Inventory management

Supply chain governance committees in Medicines and Vaccines closely monitor the inventory status and delivery of our products. Our core commercial cycle links supply chain forecasting with our commercial ambition. It is designed to reduce the risk of demand fluctuations and manage temporary shortages in supply. We periodically review each node of our supply chains to ensure we hold adequate safety stocks, whilst balancing working capital. We put particular emphasis on mitigating supply risks associated with medically-critical, high-revenue products and new product launches, for example using dual sourcing for key products or APIs. We use the monthly Performance Management Process across our supply chains to monitor business activity and highlight adverse trends in supply, operations, budget and workforce capability.

Business continuity

Crisis management and business continuity plans are in place across our supply chains, which include authorised response and recovery strategies, key areas of responsibility and clear communication routes. We regularly use business continuity plans to manage potential supply disruptions. Our manufacturing sites have crisis management plans in place. These plans are tested at least annually to ensure maintenance of core skills in crisis management.

Shareholder information

Share capital and control

Details of our issued share capital and the number of shares held in Treasury as at 31 December 2023 can be found in Note 37 to the financial statements, 'Share capital and share premium account'.

Our ordinary shares are listed on the London Stock Exchange (LSE) and are also quoted on the New York Stock Exchange (NYSE) in the form of American Depositary Shares (ADS). Each ADS represents two Ordinary Shares. For details of listed debt and where it is listed refer to Note 30 to the financial statements, 'Net debt'.

Holders of Ordinary Shares and ADS are entitled to receive dividends (when declared) and a copy of the company's Annual Report (if elected). They are also entitled to attend, speak, appoint proxies and exercise voting rights at general meetings of the company

There are no restrictions on the transfer, or limitations on the holding, of Ordinary Shares and ADS and no requirements to obtain approval prior to any transfers. No Ordinary Shares or ADS carry any special rights with regard to control of the company and there are no restrictions on voting rights. Major shareholders have the same voting rights per share as all other shareholders. There are no known arrangements under which financial rights are held by a person other than the holder of the shares and no known agreements on restrictions on share transfers or on voting rights.

Shares acquired through the Group's employee share plans rank equally with the other shares in issue and have no special rights. The trustees of our Employee Share Ownership Plan Trusts have waived their rights to dividends on shares held by those Trusts.

Demerger of Haleon and Share Consolidation

As reported previously, on 18 July 2022 the company completed the demerger of the Consumer Healthcare business from the Group. More details can be found on www.gsk.com/en-gb/haleon-cmd-to-demerger-archive. On 19 July 2022, shareholders received four new GSK plc shares of nominal value of 31^{1/4} pence each for every five GSK plc shares of nominal value of 25 pence each.

The Group reduced its share holding in Haleon plc during the course of the financial year ended 31 December 2023 to 7.4%. More information can be found in Note 22 Current Equity Investments. On 17 January 2024, the Group reduced its shareholding by 3.2%, GSK now holds approximately 385 million ordinary shares in Haleon plc representing over 4.0% of the issued share capital of Haleon. More information can be found in Note 48 Post Balance Sheet Events.

Exchange controls and other limitations affecting holders

Other than certain economic sanctions, which may be in force from time to time, there are currently no applicable laws, decrees or regulations in force in the UK restricting the import or export of capital or restricting the remittance of dividends or other payments to holders of the company's shares who are non-residents of the UK.

Similarly, other than certain economic sanctions which may be in force from time to time, there are no limitations relating only to non-residents of the UK under English law or the company's

Articles of Association on the right to be a holder of, and to vote in respect of, the company's shares.

Interests in voting rights

Other than as stated below, as far as as the company is aware, there are no persons with significant direct or indirect holdings in the company. Information provided to the company pursuant to the FCA's Disclosure Guidance and Transparency Rules (DTR 5) is published on a Regulatory Information Service and on the company's website, gsk.com.

The company has received notifications in accordance with DTR 5 of the following notifiable interests in the voting rights in the company's issued share capital:

	31 December 2023		23 February 2024	
	No. of voting rights	Percentage of total voting rights ⁽¹⁾	No. of voting rights	Percentage of total voting rights ⁽¹⁾
BlackRock, Inc.	231,975,400 ⁽²⁾	5.69 %	231,975,400 ⁽²⁾	5.69 %
Dodge & Cox	253,464,108 ⁽³⁾	5.04 %	253,464,108 ⁽³⁾	5.04 %

(1) Percentage of total voting rights at the date of notification to the company.

(2) Comprising an indirect interest in 229,134,683 Ordinary Shares and a holding of 2,840,717 Qualifying Financial Instruments (Contracts for Difference).

(3) Comprising an indirect interest in 99,377,874 Ordinary Shares and 154,086,234 ADS.

The company has not acquired or disposed of any interests in its own shares during the period under review.

Share buy-back programme

The Board has been authorised to issue and allot Ordinary Shares under Article 9 of the company's Articles of Association. The power under Article 9 and the authority for the company to make purchases of its own shares are subject to shareholder authorities which are sought on an annual basis at our Annual General Meeting (AGM). Any shares purchased by the company may be cancelled, held as Treasury shares or used for satisfying share options and grants under the Group's employee share plans.

Our programme covers purchases of shares for cancellation or to be held as Treasury shares, in accordance with the authority renewed by shareholders at the AGM in May 2023, when the company was authorised to purchase a maximum of just over 409 million shares.

In determining specific share repurchase levels, the company considers the development of free cash flow during the year. No Treasury shares have been purchased since 2014. Details of shares purchased, cancelled, held as Treasury shares and subsequently transferred from Treasury to satisfy awards under the Group's employee share plans are disclosed in Note 37 to the financial statements, 'Share capital and share premium account'. The company confirms that it does not currently intend to make any market purchases in 2024. The company will continue to review the potential for future share buy-backs in line with its usual annual cycle and subject to return and ratings criteria.

Shareholder information continued

Share capital and control continued

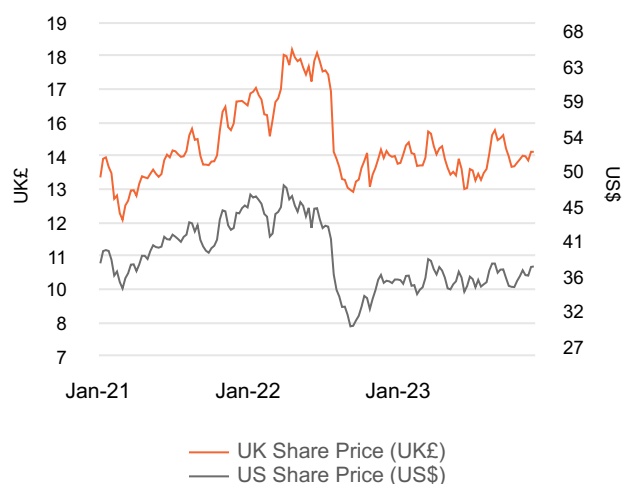
Market capitalisation

The market capitalisation, based on shares in issue excluding Treasury shares, of GSK at 31 December 2023 was £62.5 billion. At that date, GSK was the 8th largest company by market capitalisation in the FTSE index.

Share price	2023 £	2022 £	2021 £
At 1 January	14.51	16.25	13.42
At 31 December	14.50	14.38	16.07
Increase/(decrease)	(0.06)%	(12)%	20 %
High during the year	15.36	18.31	16.19
Low during the year	13.16	12.96	11.91

The table above sets out middle market closing prices. The company's share price decreased by (0.06)% in 2023. This compares with a decrease in the FTSE 100 index of 12% during the year. The middle market closing share price on 23 February 2024 was £16.72.

Share price trend in the three years ended 31 December 2023



Nature of trading market

The following table sets out, for the periods indicated, the high and low middle market closing prices for the company's Ordinary Shares on the LSE and for the ADS on the NYSE.

	Ordinary Shares		ADS	
	UK£ per share		US\$ per share	
	High	Low	High	Low
February 2024*	16.78	15.86	33.31	31.90
January 2024	15.82	14.80	40.10	37.51
December 2023	14.62	14.19	37.10	35.88
November 2023	14.26	13.82	34.17	35.99
October 2023	15.21	14.33	37.56	34.56
September 2023	13.36	13.74	38.07	34.41
Quarter ended 31 December 2023	15.21	13.82	37.56	34.17
Quarter ended 30 September 2023	15.36	13.16	38.07	33.81
Quarter ended 30 June 2023	15.23	13.46	38.32	33.60
Quarter ended 31 March 2023	15.03	13.77	36.43	33.50
Quarter ended 31 December 2022	14.92	13.20	37.92	30.00
Quarter ended 30 September 2022	18.23	12.96	44.53	28.67
Quarter ended 30 June 2022	18.31	16.72	47.70	41.98
Quarter ended 31 March 2022	17.27	15.01	47.66	40.17
Year ended 31 December 2021	16.19	13.80	44.44	38.13
Year ended 31 December 2020	14.68	12.92	39.17	33.42
Year ended 31 December 2019	18.19	14.36	47.32	37.83

* to 23 February 2024

Shareholder information continued

Analysis of shareholdings at 31 December 2023

	Number of accounts	% of total accounts	% if total shares	Number of shares
Holding of shares				
Up to 1,000	46,607	75.48	0.32	13,747,981
1,001 to 5,000	11,313	18.32	0.55	23,914,101
5,001 to 100,000	2,843	4.60	1.21	52,308,743
100,001 to 1,000,000	654	1.06	5.31	229,085,155
Over 1,000,000	328	0.53	92.60	3,993,090,003
	61,745	100.00	100.00	4,312,145,983
Held by				
Institutional and corporate holders	2,153	3.49	61.86	2,667,435,551
Individuals and other corporate bodies	59,590	96.51	13.86	597,606,148
Guaranty Nominees Limited (ADR programme)	1	0.00	19.71	850,036,115
Held as Treasury shares by GSK	1	0.00	4.57	197,068,169
	61,745	100.00	100.00	4,312,145,983

JP Morgan Chase Bank NA is the Depository for the company's American Depositary Receipt (ADR) programme. The company's ADS are listed on the NYSE. Ordinary Shares representing the company's ADR programme, which is managed by the Depository, are registered in the name of Guaranty Nominees Limited. At 23 February 2024, Guaranty Nominees Limited held 832,929,801 Ordinary Shares representing 20.23% of the issued share capital (excluding Treasury shares).

At 23 February 2024, the number of holders of Ordinary Shares in the US was 842 with holdings of 689,588 Ordinary Shares, and the number of registered holders of ADS was 15,511 with holdings of 416,464,900 ADS. Certain of these Ordinary Shares and ADS were held by brokers or other nominees. As a result, the number of holders of record or registered holders in the US is not representative of the number of beneficial holders or of the residence of beneficial holders.

Dividends

The company pays dividends quarterly and continues to return cash to shareholders through its dividend policy. Dividends remain an essential component of total shareholder return and GSK recognises the importance of dividends to shareholders.

On 23 June 2021, at the GSK Investor Update, GSK set out that from 2022 a progressive dividend policy will be implemented guided by a 40 to 60 percent pay-out ratio through the investment cycle. The dividend policy, the total expected cash distribution, and the respective dividend pay-out ratios for GSK remain unchanged.

Dividends per share

The table below sets out the dividend per share and per ADS for the last five years. The dividend per ADS is translated into US dollars at applicable exchange rates.

Year	pence	US\$ ⁽¹⁾
2023	58 ⁽²⁾	— ⁽⁴⁾
2022	61.25 ⁽³⁾	2.00
2021	80	2.16
2020	80	2.12
2019	80	2.01

(1) An annual fee of \$0.03 per ADS (or \$0.0075 per ADS per quarter) will be charged by the Depository. The amounts shown are the dividends paid per ADS before the annual fee is charged.

(2) Dividends declared and paid in respect of 2023 were 14p per share for Q1 2023, 14p per share for Q2 2023 and 14p per share for Q3 2023. A dividend of 16p per share has been declared for Q4 2023.

(3) Adjusted for the Share Consolidation (2022 only; prior years have not been adjusted).

(4) The Q4 2023 ordinary dividend receivable by ADS holders will be calculated based on the exchange rate on 9 April 2024. The cumulative dividend receivable by ADS holders for Q1, Q2 and Q3 2023 was £1.06.

The expected dividend for 2024 is 60p per Ordinary Share.

Details of the dividends declared, the amounts and the payment dates are given in Note 16 to the financial statements, 'Dividends'.

2024 Dividend calendar

Quarter	Ex-dividend date	ADS Ex-dividend date	Record date	Payment date
Q4 2023	22 February 2024	22 February 2024	23 February 2024	11 April 2024
Q1 2024	16 May 2024	16 May 2024	17 May 2024	11 July 2024
Q2 2024	15 August 2024	16 August 2024	16 August 2024	10 October 2024
Q3 2024	November 2024	November 2024	November 2024	9 January 2025
Q4 2024	20 February 2025	21 February 2025	21 February 2025	10 April 2025

Shareholder information continued

Financial calendar 2024

Event	Date
Quarter 1 results announcement	1 May 2024
Annual General Meeting	8 May 2024
Quarter 2 results announcement	31 July 2024
Quarter 3 results announcement	30 October 2024
Preliminary/Quarter 4 Results announcement	5 February 2025
Annual Report publication	February/March 2024
Annual Report distribution	March 2024

Information about the company, including the share and ADS price, is available on our website at gsk.com. Information made available on the website does not constitute part of this Annual Report.

Stock Exchange announcement notifications

We provide shareholders with a service to receive automatic email notifications when we publish a stock exchange announcement. To receive email notifications, please sign up for announcements at gsk.com in the Investors section.

Results announcements

Results announcements are issued to the LSE and are available on its news service. They are also sent to the US Securities and Exchange Commission (SEC) and the NYSE, issued to the media and made available on our website.

Financial reports

The company publishes an Annual Report which is made available on our website from the date of publication. Shareholders may elect to receive notification by email of the publication of Annual Reports by registering on www.shareview.co.uk, and may also elect to receive a printed copy of the Annual Report by contacting our registrar, Equiniti Limited.

Copies of previous Annual Reports are available on our website. Printed copies can also be obtained from our registrar (see page 301 for the contact details).

Annual General Meeting 2024

Our Annual General Meeting (AGM) will be held at 2.30pm (UK time) on Wednesday, 8 May 2024 at the Royal Lancaster London, Lancaster Terrace, London W2 2TY and will also be broadcast live for you to join electronically.

The AGM is the company's principal forum for communication with private shareholders. In addition to the formal AGM business, there will be a presentation by the CEO on the performance of the Group and its future development. There will be an opportunity for questions to be asked of the Board and Chairs of the Board's Committees will be available to take questions relating to their roles.

Further details on how to access the AGM electronically or attend in person, ask questions and vote, can be found in the notice of Annual General Meeting 2024 (AGM Notice) which will be made available on our website at gsk.com on or around 25 March 2024.

Investors holding shares through a nominee service should arrange with that service for them to be appointed as a proxy in respect of their shareholding to attend and vote at the meeting electronically.

ADS holders wishing to attend the meeting electronically should refer to the AGM Notice for details on how to request a proxy appointment from the Depositary, JP Morgan Chase Bank NA. This will enable them to attend, ask questions and vote electronically on the business to be transacted at the meeting.

ADS holders are reminded that if they do not instruct the Depositary as to the way in which the shares represented by their ADS should be voted by completing and returning the voting card provided by the Depositary, their shares will not be voted.

Documents on display

The Articles of Association of the company and Directors' service contracts or, where applicable, letters of appointment between Directors and the company or any of its subsidiaries (and any side letters relating to severance terms and pension arrangements) are available for inspection at the company's registered office and will be made available for inspection at the AGM.

Shareholder information continued

Tax information for shareholders

A summary of certain UK tax and US federal income tax consequences for holders of shares and ADS who are citizens of the UK or the US is set out below. It is not a complete analysis of all the possible tax consequences of the purchase, ownership or sale of these securities. It is intended only as a general guide. Holders are advised to consult their advisers with respect to the tax consequences of the purchase, ownership or sale of their shares or ADS and the consequences under state and local tax laws in the US and the implications of the current UK/US tax conventions.

US holders of ADS generally will be treated as the owners of the underlying shares for the purposes of the current UK/US double taxation conventions relating to income and gains (Income Tax Convention), estate and gift taxes (Estate and Gift Tax Convention), and for the purposes of the Internal Revenue Code of 1986, as amended.

UK shareholders

This summary only applies to a UK resident shareholder that holds shares as capital assets.

Taxation of dividends

For the 2023/24 UK tax year, UK resident individuals are entitled to a dividend tax allowance of up to £1,000, so that the first £1,000 of dividends received in a tax year will be free of tax. Dividends in excess of this allowance will be taxed at 8.75% for basic rate taxpayers, 33.75% for higher rate taxpayers and 39.35% for additional rate taxpayers. Note that from 6 April 2024 the dividend allowance will be reduced to £500.

UK resident shareholders that are corporation taxpayers should note that dividends payable on ordinary shares are generally entitled to exemption from corporation tax.

Taxation of capital gains

UK resident shareholders may be liable for UK tax on gains on the disposal of shares or ADS.

For disposals by individuals in the 2023/24 UK tax year, a taxable capital gain accruing on a disposal of shares or ADS will be taxed at 10% for basic rate taxpayers, or 20% if, after all allowable deductions, the individual's taxable income for the year exceeds the basic rate income tax banding. Note this is following the use of any exemptions available to the individual taxpayer such as the annual exempt amount.

Corporation taxpayers may be entitled to an indexation allowance which applies to reduce capital gains to the extent that such gains arise due to inflation. Indexation allowance may reduce a chargeable gain but will not create an allowable loss. For assets acquired on or before 1 January 2018, legislation in the Finance Act 2018 freezes the level of indexation allowance that is given in calculating a company's chargeable gains at the value that would apply to the disposal of an asset in December 2017. For assets acquired from 1 January 2018 onwards, legislation in the Finance Act 2018 removes any indexation allowance on disposal.

Inheritance tax

Individual (UK-domiciled or otherwise) shareholders may be liable to UK inheritance tax on the transfer of shares or ADS. Exposure to a UK inheritance tax charge typically occurs on the death of the asset owner. However, transfers of shares (other than commercial sales) within seven years of death remain relevant to any inheritance tax exposure at death. Further, transfers to a trust arrangement during lifetime can give rise to an immediate inheritance tax charge.

Tax may be charged on the amount by which the value of the shareholder's estate is reduced as a result of any transfer by way of lifetime gift or other disposal at less than full market value. In the case of a bequest on death, tax may be charged on the value of the shares at the date of the shareholder's death. Where an exposure to UK inheritance tax and US estate or gift tax exists, careful planning must be undertaken to understand the opportunity to utilise the US/UK Estate and Gift Double Tax Convention to manage tax credits and avoid double taxation.

The overall exposure will be dependent on the specific circumstances of each situation and it is also important to note that tax charges may arise in other jurisdictions. Bespoke advice tailored to an individual's personal circumstances should therefore be obtained from a tax professional.

Stamp duty and stamp duty reserve tax

UK stamp duty and/or stamp duty reserve tax (SDRT) will, subject to certain exemptions, be payable on the transfer of shares at a rate of 0.5% (rounded up to the nearest £5 in the case of stamp duty) of the consideration for the transfer. Notwithstanding this, provided that an instrument is executed in pursuance of the agreement that gave rise to the charge to SDRT and that instrument is stamped within six years of the agreement (including being stamped as exempt) any SDRT charge should be cancelled and any SDRT which has already been paid will be repaid. Where listed shares are transferred to a company connected to the transferor the chargeable consideration will be deemed to be not less than the market value of the shares transferred. This market value override also applies where non-listed shares are transferred to a company connected to the transferor where the consideration includes an issue of shares.

US shareholders

This summary only applies to a shareholder (who is a citizen or resident of the US or a domestic corporation or a person that is otherwise subject to US federal income tax on a net income basis in respect of the shares or ADS) that holds shares or ADS as capital assets, is not resident in the UK for UK tax purposes and does not hold shares for the purposes of a trade, profession or vocation that is carried on in the UK through a branch or agency.

The summary also does not address the tax treatment of holders that are subject to special tax rules, such as banks, tax-exempt entities, insurance companies, dealers in securities or currencies, persons that hold shares or ADS as part of an integrated investment (including a 'straddle') comprised of a share or ADS and one or more other positions, and persons that own (directly, indirectly or constructively) 10% or more of the company's stock (by vote or value), nor does it address tax treatment that may be applicable as a result of international income tax treaties.

Shareholder information continued

Tax information for shareholders continued

Taxation of dividends

The gross amount of dividends received is treated as foreign source dividend income for US tax purposes. It is not eligible for the dividend received deduction allowed to US corporations. Dividends on ADS are payable in US dollars; dividends on Ordinary Shares are payable in sterling. Dividends paid in sterling will be included in income in the US dollar amount calculated by reference to the exchange rate on the day the dividends are received by the holder. Subject to certain exceptions for short-term or hedged positions, an individual eligible US holder will be subject to US taxation at a maximum federal rate of 23.8% plus applicable state and local tax in respect of qualified dividends. A qualified dividend as defined by the US Internal Revenue Service (IRS) is a dividend that meets the following criteria:

1. It must be issued by a US corporation, a corporation incorporated in a US possession, or a corporation that is eligible for the benefits of a comprehensive income tax treaty deemed satisfactory, as published by the IRS.
2. The dividends are not of a type listed by the IRS as dividends that do not qualify.
3. The required dividend holding period has been met. The shares must have been owned by you for more than 60 days of the 'holding period' – which is defined as the 121-day period that begins 60 days before the ex-dividend date, or the day in which the stock trades without the dividend priced in. For example, if a stock's ex-dividend date is 1 October, the shares must be held for more than 60 days in the period between 2 August and 30 November of that year in order to count as a qualified dividend.

Dividends that are not qualified are subject to taxation at the US federal graduated tax rates, at a maximum rate of 40.8%. Some types of dividends are automatically excluded from being qualified dividends, even if they meet the other requirements. These include (but are not limited to):

- Capital gains distributions
- Dividends on bank deposits
- Dividends held by a corporation in an Employee Stock Ownership Plan (ESOP)
- Dividends paid by tax-exempt corporations.

US state and local tax rates on qualified and non-qualified dividends may vary and would be assessed in addition to the federal tax rates communicated above.

Taxation of capital gains

Generally, US holders will not be subject to UK capital gains tax, but will be subject to US tax on capital gains realised on the sale or other disposal of shares or ADS. Such gains will be long-term capital gains (subject to reduced rates of taxation for individual holders) if the shares or ADS were held for more than one year, from the date the shares were vested/released. Short-term capital gains can be subject to taxation of rates of up to 40.8%, whereas long-term capital gains may be subject to rates of up to 23.8%. State and local tax rates on capital gains may also apply.

Information reporting and backup withholding

Dividends and payments of the proceeds on a sale of shares or ADS, paid within the US or through certain US-related financial intermediaries, are subject to information reporting and may be subject to backup withholding unless the US holder is a corporation or other exempt recipient or provides a taxpayer identification number and certifies that no loss of exemption has occurred. Non-US holders generally are not subject to information reporting or backup withholding, but may be required to provide a certification of their non-US status in connection with payments received. Any amounts withheld will be allowed as a refund or credit against a holder's US federal income tax liability provided the required information is furnished to the IRS.

Estate and gift taxes

Under the Estate and Gift Tax Convention, a US shareholder is not generally subject to UK inheritance tax. However, a US holder may be subject to US federal estate and gift tax.

Stamp duty

UK stamp duty and/or SDRT will, subject to certain exemptions, be payable on any transfer of shares to the ADS custodian or depository at a rate of 1.5% of the amount of any consideration provided (if transferred on sale), or their value (if transferred for no consideration).

However, no stamp duty or SDRT should be payable on the transfer of, or agreement to transfer an ADS or on transfers within the clearance service. Notwithstanding the above, where the clearance service operator has made an election under s97A Finance Act 1986, broadly the 1.5% stamp duty/SDRT charge should not arise on the transfer into the clearance service, but transfers to, and within, the system (where there is a change in beneficial ownership) would attract a 0.5% charge.

Demerger and share consolidation

A summary of certain UK and US tax consequences in respect of the demerger of Haleon plc and the consolidation of the company's share capital, relevant to the company's shareholders who are resident (or, in the case of individuals, resident and domiciled) in the UK for UK tax purposes or who are citizens of or resident in the US for US tax purposes, is set out in Part 6 of the circular in relation to the Demerger and the Share Consolidation published on 1 June 2022 (Circular) (pages 83 to 89). The Circular, along with other information regarding the demerger and share consolidation can be found at gsk.com in the demerger section.

Further information on the tax base cost allocation to assist UK shareholders apportion their base cost between their GSK plc shares and Haleon plc shares for UK capital gains tax purposes following the demerger, including a worked example, can be found in the Tax section at gsk.com in the demerger section.

Other statutory disclosures

Shareholder services and contacts

Registrar

The company's registrar is:

Equiniti Limited
Aspect House, Spencer Road, Lancing, BN99 6DA
www.shareview.co.uk
Tel: +44 (0)371 384 2991*

Equiniti provides a range of services for shareholders:

Service	What it offers	How to participate
Dividend Reinvestment Plan (DRIP)	As an alternative to receiving cash dividends you may choose to reinvest your dividends to buy more GSK shares.	A DRIP election form, Terms and Conditions and information on fees can be downloaded from www.shareview.co.uk or requested by contacting Equiniti.
Dividend payment direct to your bank account (bank mandate)	All dividends are paid directly into your bank or building society account. To receive your cash dividends, you must provide Equiniti with your bank or building society account details. This is a quick and secure method of payment.	A dividend bank mandate form can be downloaded from www.shareview.co.uk or requested by contacting Equiniti.
Dividend payment direct to bank account for overseas shareholders (Overseas Payment Service)	Equiniti can convert your dividend into your local currency and send it direct to your local bank account. The Overseas Payment Service is available in approximately 100 countries worldwide.	More information on the Overseas Payment Service (including information on fees) can be found at www.shareview.co.uk or by contacting Equiniti.
Electronic communications	Shareholders may elect to receive electronic notifications of company communications including our Annual Report, dividend payments, dividend confirmations and the availability of online voting for all general meetings. Each time GSK publishes shareholder documents you will receive an email containing a link to the document or relevant website.	Please register at www.shareview.co.uk .
Shareview portfolio service	This enables you to create a free online portfolio to view your share balance and movements, update your address and dividend payment instructions and register your votes for our general meetings.	Please register at www.shareview.co.uk .
Deduplication of publications or mailings	If you receive duplicate copies of mailings, you may have more than one account. Please contact Equiniti and they will arrange for your accounts to be merged into one for your convenience and to avoid waste and unnecessary costs.	Please contact Equiniti.
Share dealing service[†] (please note that market trading hours are from 8.00am to 4.30pm UK time, Monday to Friday (excluding public holidays in England and Wales))	Shareholders may trade shares, either held in certificated form or in our Corporate Sponsored Nominee, online, by telephone or via postal dealing service provided by Equiniti Financial Services Limited.	More information on the share dealing service (including information on fees) can be found at www.shareview.co.uk/dealing For online transactions, please log on to: www.shareview.co.uk/dealing . For telephone transactions, please call: 0345 603 7037 (in the UK) or +44 (0)345 603 7037 (outside the UK). Lines are open from 8.00am to 4.30pm UK time, Monday to Friday (excluding UK public holidays). For postal transactions, please call: 0371 384 2991* to request a dealing form.
Corporate Sponsored Nominee Account	This is a convenient way to manage your shares without requiring a share certificate. The service provides a facility for you to hold your shares in a nominee account sponsored by the company. You will continue to receive dividend payments and can attend and vote at the company's general meetings. Shareholders' names do not appear on the publicly available share register and the service is free to join.	An application form can be requested from www.shareview.co.uk or by contacting Equiniti.
Individual Savings Accounts (ISAs)[†]	Equiniti Financial Services Limited provide the EQi Flexible ISA to hold GSK shares.	Details (including information on fees) are available from www.eqi.co.uk or can be requested by calling the Equiniti Customer Experience Team on 0345 0700 720. Lines are open 8:00am to 5:30pm, UK time Monday to Friday (excluding UK public holidays).

* Lines are open from 8.30am to 5.30pm, UK time Monday to Friday (excluding public holidays in England and Wales). Please use the country code when dialling from outside the UK.

† The provision of share dealing details is not intended to be an invitation or inducement to engage in an investment activity. Advice on share dealing should be obtained from a stockbroker or independent financial adviser.

Other statutory disclosures continued

Shareholder services and contacts continued

ADS Depository

The ADR programme is administered by JPMorgan Chase Bank, N.A.:

Regular Correspondence:
EQ Shareowner Services
P.O. Box 64504
St. Paul, MN 55164-0504

Delivery of Stock Certificates and Overnight Mail:
EQ Shareowner Services
1110 Centre Pointe Curve, Suite 101
Mendota Heights, MN 55120-4100

shareowneronline.com/informational/contact-us/
From the US: +1 877 353 1154
From outside the US: +1 651 453 2128

The Depository also provides Global Invest Direct, a direct ADS purchase/sale and dividend reinvestment plan for ADS holders. For details on how to enrol, please visit www.adr.com or call the above helpline number to obtain an enrolment pack.

Contacts

Investor relations

Investor relations may be contacted as follows:

UK

980 Great West Road
Brentford, Middlesex, TW8 9GS
Tel: +44 (0)20 8047 5000

US

2929 Walnut Street
Philadelphia PA 19104
Tel: +1 888 825 5249 (US toll free)
Tel: +1 215 751 4000 (outside the US)-

GSK Response Center

Tel: +1 888 825 5249 (US toll free)
Tel: +1 215 751 4600 (outside the US)

Share scam alert

If you receive an unsolicited telephone call offering to sell or buy your shares, please take extra care. The caller may be part of a highly organised financial scam.

If you are a UK shareholder, please contact the Financial Conduct Authority at www.fca.org.uk/consumers or on its consumer helpline:

Tel: 0800 111 6768 (in the UK)*

Tel: +44 207 066 1000 (outside the UK)*

* Lines are open from 8.00am to 6.00pm, UK time, Monday to Friday, except UK public holidays, and 9.00am to 1.00pm on Saturdays.

Donating shares to Save the Children

In 2013, GSK embarked on an ambitious global partnership with Save the Children to share our expertise and resources with the aim of finding innovative ways to reduce the number of children dying from preventable diseases.

Shareholders with a small number of shares, the value of which makes it uneconomical to sell, may wish to consider donating them to Save the Children. Donated shares will be aggregated and sold on behalf of Save the Children who will use the funds raised to help them reach the above goal.[†]

To obtain a share donation form, please contact our registrar, Equiniti, which is managing the donation and sale of UK shares to Save the Children free of charge.

[†] The provision of share dealing details is not intended to be an invitation or inducement to engage in an investment activity. Advice on share dealing should be obtained from a stockbroker or independent financial adviser.

Other statutory disclosures continued

US law and regulation

A number of provisions of US law and regulation apply to the company because our shares are quoted on the NYSE in the form of ADS.

NYSE rules

In general, the NYSE rules permit the company to follow UK corporate governance practices instead of those applied in the US, provided that we explain any significant variations. This explanation is contained in our Form 20-F, which can be accessed from the SEC's EDGAR database or via our website. NYSE rules require us to file annual and interim written affirmations concerning our Audit & Risk Committee (ARC) and our statement on significant differences in corporate governance.

Sarbanes-Oxley Act of 2002

Following a number of corporate and accounting scandals in the US, 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 SEC, the company has an established Disclosure Committee. The Committee reports to the CEO, the CFO and to the ARC. It is chaired by the Company Secretary and its members consist of senior managers from finance, legal, corporate communications and investor relations.

Where appropriate, external legal counsel, the external auditors, our sponsor bank, and internal experts are invited to attend the Disclosure Committee's meetings periodically. The Committee 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 Annual Report and the Annual Report on Form 20-F. In 2023, the Committee met 17 times, including for the purpose of receiving relevant and appropriate training.

Sarbanes-Oxley requires that the Annual Report on Form 20-F contains a statement as to whether a member of the ARC is an audit committee financial expert, as defined in rules under Sarbanes-Oxley. Such a statement for the relevant members of the ARC (Charles Bancroft) is included in the Board Committee information area of the Corporate Governance report on page 118 and in his biography on page 109.

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 requires the CEO and the CFO to complete formal certifications, confirming that:

- they have each reviewed the Annual Report on Form 20-F;
- based on their knowledge, the Annual Report on Form 20-F contains 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 Annual Report on 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 Annual Report on 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 Annual Report on Form 20-F any changes in internal controls over financial reporting during the period covered by the Annual Report on 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 auditor 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 2023.

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.

Other statutory disclosures continued

US law and regulation continued

The CEO and CFO expect to complete these certifications and report their conclusions on the effectiveness of disclosure controls and procedures in March 2024, following which the certifications will be filed with the SEC as part of our Group's Annual Report on Form 20-F.

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, as amended (the Exchange Act)):

- 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, Internal Control – Integrated Framework (2013) issued by the Committee of Sponsoring Organisations of the Treadway Commission (COSO).
- There have been no changes in the Group's internal control over financial reporting during 2023 that have materially affected, or are reasonably likely to materially affect, the Group's internal control over financial reporting.
- Management has assessed the effectiveness of internal control over financial reporting as at 31 December 2023 and its conclusion will be filed as part of the Group's Annual Report on Form 20-F.
- Deloitte LLP, which has audited the consolidated financial statements of the Group for the year ended 31 December 2023, has also assessed the effectiveness of the Group's internal control over financial reporting under Auditing Standard 2201 of the Public Company Accounting Oversight Board (United States). Their audit report will be filed with the Group's Form 20-F.

Section 13(r) of the Exchange Act

Section 13(r) of the Exchange Act requires issuers to make specific disclosure in their annual reports of certain types of dealings with Iran, including transactions or dealings with government-owned or-controlled entities, as well as dealings with entities sanctioned for activities related to terrorism or proliferation of weapons of mass destruction, even when those activities are not prohibited by US law and do not involve US persons.

The Group exports certain medicines to Iran, via sales by non-US entities that are not subsidiaries of a US entity to a distributor in Iran pursuant to a specific licence issued by the Office of Foreign Assets Control.

The Group does not regularly receive information regarding the identity of the distributor's downstream customers and intermediaries in Iran, and it is possible that these parties include entities, such as hospitals and pharmacies, that are owned directly or indirectly by the Iranian government or by persons or entities sanctioned in connection with terrorism or proliferation activities.

As the Group does not regularly receive information regarding the identity of its distributor's downstream customers and intermediaries it cannot establish the proportion of gross revenue or sales potentially attributable to entities affiliated with the Iranian government or parties sanctioned for disclosable activities. As a result, the Group is reporting the entire gross revenues (£16.89 million) and net profits (£8.42 million) from the Group's sales to Iran in 2023.

Some hospitals or other medical facilities in Lebanon may be affiliated with or controlled by Hezbollah or other groups that are designated by the United States pursuant to Executive Order 13224. Again, the Group does not deal directly with such hospitals or facilities and instead sells through distributors. The Group is unable to establish the proportion of gross revenue or sales potentially attributable to reportable activities. As a result, the Group is reporting the entire gross revenues (£6.02 million) and net losses (£4.2 million) from the Group's sales to Lebanon in 2023.

In addition to Section 13(r) of the Exchange Act, US law generally restricts dealings by US persons and dealings that otherwise are subject to US jurisdiction with certain countries or territories that are subject to comprehensive sanctions, currently Crimea, Cuba, the so-called Donetsk People's Republic, Iran, the so-called Luhansk People's Republic, North Korea and Syria, as well as with the Government of Venezuela (though not with the country of Venezuela as a whole). The Group engages in some activity in certain such jurisdictions having assessed applicable licences and exemptions

While we believe the Group complies with all applicable US sanctions in all material respects, such laws are complex and continue to evolve rapidly.

Other statutory disclosures continued

Donations to political organisations and political expenditure

To ensure a consistent approach to political contributions across the Group, in 2009 a global policy was introduced to voluntarily stop all corporate political contributions.

In the period from 1 January 2009 to 31 December 2023, the Group has not made any political donations to EU or non-EU organisations.

Notwithstanding the introduction of this policy, in accordance with the Federal Election Campaign Act in the US, we continue to support an employee-operated Political Action Committee (PAC) that facilitates voluntary political donations by eligible GSK employees.

The PAC is not controlled by GSK. Decisions on the amounts and recipients of contributions are governed by the PAC Board of Directors. Contributions to the PAC are made by participating eligible employees exercising their legal right to pool their resources and make political contributions, which are subject to strict limitations under US law. In 2023, a total of US\$325,750 (2022: US\$360,950) was donated to political organisations by the GSK employee PAC.

English law requires prior shareholder approval for political contributions to EU political parties and independent election candidates as well as for any EU political expenditure. The definitions of political donations, political expenditure and political organisations used in the legislation are, however, quite broad. In particular, the definition of EU political organisations may extend to bodies such as those concerned with policy review, law reform, the representation of the business community and special interest groups such as those concerned with the environment, which the company and its subsidiaries might wish to support.

As a result, the definitions may cover legitimate business activities not in the ordinary sense considered to be political donations or political expenditure, nor are they designed to support any political party or independent election candidate.

Therefore, notwithstanding our policy, and while we do not intend to make donations to any EU political parties or organisations, nor to incur any EU political expenditure, we annually seek shareholder authorisation for any inadvertent expenditure.

The authority is a precautionary measure to ensure that the company and its subsidiaries do not inadvertently breach the legislation.

This authorisation process, for expenditure of up to £100,000 each year, dates back to the AGM held in May 2001, following the introduction of the Political Parties, Elections and Referendums Act 2000. The authority has since been renewed annually.

Other statutory disclosures continued

Group companies

In accordance with Section 409 of the Companies Act 2006 a full list of subsidiaries, associates, joint ventures and joint arrangements, the address of the registered office and effective percentage of equity owned, as at 31 December 2023 are disclosed below. Unless otherwise stated the share capital disclosed comprises ordinary shares which are indirectly held by GSK plc. The percentage held by class of share is stated where this is less than 100%. Unless otherwise stated, all subsidiary companies have their registered office and are tax resident in their country of incorporation.

Name	Security	Registered address
Wholly owned subsidiaries		
14245563 Canada Inc.	Common	275 Armand-Frappier Boulevard, Laval ON H7V 4A7, Canada
14934792 Canada Inc.	Common	100 Milverton Drive, Suite 800, Mississauga ON L5R 4H1, Canada
1506369 Alberta ULC	Common	3500 855-2nd Street SW, Calgary AB T2P 4J8, Canada
Action Potential Venture Capital Limited	Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
Adechsa GmbH (ii)	Ordinary	c/o PRV Provides Treuhandgesellschaft AG, Dorfstrasse 38, 6341, Baar, Switzerland
Affinivax, Inc.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Allen & Hanburys Limited (ii)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Allen & Hanburys Pharmaceutical Nigeria Limited	Ordinary	49, Town Planning Way, Ilupeju, Lagos, Nigeria
Allen Pharmazeutika Gesellschaft m.b.H.	Ordinary	Wienerbergstraße 7, Wien, 1100, Austria, Austria
Beecham Group p.l.c	£0.05 Ordinary B; £0.20 Ordinary A	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Beecham Pharmaceuticals (Pte) Limited	Ordinary	38 Quality Road, Jurong Industrial Estate, Jurong, 618809, Singapore
Beecham Portuguesa- Produtos Farmaceuticos e Quimicos, LDA	Quota	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
Beecham S.A.	Ordinary	Avenue Fleming 20, 1300 Wavre, Belgium
Bellus Health Corp.	Common	The Corporation Trust Company, Corporation Trust Center, 1209 Orange Street, Wilmington DE 19801, United States
Bellus Health Inc	Common	275 Boulevard Armand Frappier, Laval QC H7V 4A7, Canada
Biovesta İlaçları Ltd. Sti. (ii)	Nominative	Büyükdere Caddesi No. 173, 1.Levent Plaza B Blok, 1.Levent, Istanbul, 34394, Turkey
Cascan GmbH & Co. KG	Partnership Capital	Prinzregentenplatz 9, 81675, Munich, Bavaria, Germany
Cellzome GmbH	Ordinary	Meyerhofstrasse 1, 69117, Heidelberg, Germany
Clarges Pharmaceutical Trustees Limited (ii)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Colleen Corporation	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Corixa Corporation	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Dealcyber Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Desarrollo Energia Solar Alternativa S.L.	Ordinary	Severo Ochoa, 2, Parque Tecnológico de Madrid, Tres Cantos, 28760, Madrid, Spain
Duncan Pharmaceuticals Philippines Inc.	Common	23rd Floor, The Finance Centre, 26th Street Corner 9th Avenue, Bonifacio Global City, Taguig City, 1634, Philippines
Etex Farmaceutica Ltda	Social Capital	Av. Andrés Bello 2457, Costanera Center, Torre 2, Piso 20, Providencia, Santiago, 7510689, Chile
Glaxo Group Limited	Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
Glaxo Kabushiki Kaisha (ii)	Ordinary	1-8-1 Akasaka Minato-ku, Tokyo, Japan
Glaxo New Zealand Pension Plan Trustee Limited	Ordinary	Level 2 E.2, Generator at GridAKL, 12 Madden Street, Wynyard Quarter, Auckland, 1010, New Zealand
Glaxo Operations UK Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Glaxo Properties BV	Ordinary	Van Asch van Wijkstraat 55h, 3811 LP, Amersfoort, Netherlands
Glaxo Saudi Arabia Limited	Ordinary	PO Box 22617, Area No 56 to 73, Warehouse City, First Stage Al Khomrah, Jeddah 21416, Saudi Arabia
Glaxo Verwaltungs GmbH	Ordinary	Prinzregentenplatz 9, 81675, Munich, Bavaria, Germany
Glaxo Wellcome Farmaceutica, Limitada	Ordinary	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
Glaxo Wellcome Manufacturing Pte Ltd	Ordinary	1 Pioneer Sector 1, Jurong Industrial Estate, Jurong, 628413, Singapore
Glaxo Wellcome Production	Ordinary	23 rue François Jacob, 92500, Rueil-Malmaison, France

Other statutory disclosures continued

Group companies continued

Name	Security	Registered address
Wholly owned subsidiaries continued		
Glaxo Wellcome Vidhyasom Limited (in liquidation) (ii)	Ordinary	12th Floor Wave Place, 55 Wireless Road, Lumpini, Pathumwan, Bangkok, 10 330, Thailand
Glaxo Wellcome, S.A.	Ordinary	Poligono Industrial Allendeduero, Avenida de Extremadura, 3, Aranda de Duero, 09400, Burgos, Spain
Glaxo, S.A.	Ordinary	Severo Ochoa, 2, Parque Tecnológico de Madrid, Tres Cantos, 28760, Madrid, Spain
Glaxochem Pte Ltd (iii)	Ordinary	23 Rochester Park, 139234, Singapore
GlaxoSmithKline - Produtos Farmaceuticos, Limitada	Ordinary Quota	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
GlaxoSmithKline (Cambodia) Co., Ltd.	Ordinary	5th Floor DKSH Building, No.797 Preah Monivong Boulevard (Co, Sangkat Phsar Deum Thakov, Khan Chamkarmon, Phnom Penh, Cambodia
GlaxoSmithKline (China) Investment Co Ltd	Ordinary	Room 901, 902, 903, 905, 908, 909 and 910, Unit 901, Floor 9, No. 56 Mid 4th East Ring Road, Chaoyang District, Beijing, China
GlaxoSmithKline (China) R&D Company Limited	Equity	Fl-3, No.18 Building, 999 Huanke Road, Pilot Free Trade Zone, Shanghai, 201 210, China
GlaxoSmithKline (GSK) S.R.L.	Ordinary	Str. Dr. Nicolae D. Staicovici nr. 2, Opera Center II, etaj 4, sector 5, București, Romania, 050556
GlaxoSmithKline (Ireland) Limited	Ordinary	12 Riverwalk, Citywest Business Campus, Dublin 24, Ireland
GlaxoSmithKline (Israel) Ltd	Ordinary	25 Basel Street, PO Box 10283, Petach-Tikva, 49002, Israel
GlaxoSmithKline (Private) Limited (ii)	Ordinary	Unit 3, 20 Anthony Road, Msasa, Harare, Zimbabwe
GlaxoSmithKline (Thailand) Limited	Ordinary	12th Floor Wave Place, 55 Wireless Road, Lumpini, Pathumwan, Bangkok, 10 330, Thailand
GlaxoSmithKline AB	Ordinary	Hemvarnsg. 9, 171 54, Solna, Sweden
GlaxoSmithKline AG	Ordinary	Talstrasse 3, 3053 Muenchenbuchsee, Switzerland
GlaxoSmithKline Angola Unipessoal Limitada	Quota	Luanda, Bairro Petrangol, Estrada de Cacuaco n ° 288, Angola
GlaxoSmithKline Argentina S.A.	Ordinary	Tucumán 1, piso 4, Buenos Aires, C1049AAA, Argentina
GlaxoSmithKline AS	Ordinary	Drammensveien 288, Oslo, NO-0283, Norway
GlaxoSmithKline Australia Pty Ltd	Ordinary	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
GlaxoSmithKline B.V.	Ordinary	Van Asch van, Wijkstraat 55h, 3811 LP Amersfoort, The Netherlands, Netherlands
GlaxoSmithKline Beteiligungs GmbH	Ordinary	Prinzregentenplatz 9, 81675, Munchen, Germany
GlaxoSmithKline Biologicals Kft.	Ordinary	2100 Gödöllő, Homoki Nagy István utca 1, Hungary
GlaxoSmithKline Biologicals S.A.S.	Ordinary	637 Rue des Aulnois, Saint-Amand Les Eaux, 59230, France
GlaxoSmithKline Biologicals SA	Ordinary: Preference	Rue de l'Institut 89 B-1330 Rixensart, Belgium
GlaxoSmithKline Brasil Limitada	Quotas	Estrada dos Banderiantes, 8464, Rio de Janeiro, 22783-110, Brazil
GlaxoSmithKline Capital Inc.	Common	Wilmington Trust SP Services, Inc., 1100 N. Market Street, 4th Floor, Wilmington DE 19890, United States
GlaxoSmithKline (China) R&D Company Limited	Equity	Fl-3, No.18 Building, 999 Huanke Road, Pilot Free Trade Zone, Shanghai, 201 210, China
GlaxoSmithKline (GSK) S.R.L.	Ordinary	Str. Dr. Nicolae D. Staicovici nr. 2, Opera Center II, etaj 4, sector 5, București, Romania, 050556
GlaxoSmithKline (Ireland) Limited	Ordinary	12 Riverwalk, Citywest Business Campus, Dublin 24, Ireland
GlaxoSmithKline (Israel) Ltd	Ordinary	25 Basel Street, PO Box 10283, Petach-Tikva, 49002, Israel
GlaxoSmithKline (Private) Limited (ii)	Ordinary	Unit 3, 20 Anthony Road, Msasa, Harare, Zimbabwe
GlaxoSmithKline (Thailand) Limited	Ordinary	12th Floor Wave Place, 55 Wireless Road, Lumpini, Pathumwan, Bangkok, 10 330, Thailand
GlaxoSmithKline AB	Ordinary	Hemvarnsg. 9, 171 54, Solna, Sweden
GlaxoSmithKline AG	Ordinary	Talstrasse 3, 3053 Muenchenbuchsee, Switzerland
GlaxoSmithKline Angola Unipessoal Limitada	Quota	Luanda, Bairro Petrangol, Estrada de Cacuaco n ° 288, Angola
GlaxoSmithKline Argentina S.A.	Ordinary	Tucumán 1, piso 4, Buenos Aires, C1049AAA, Argentina
GlaxoSmithKline AS	Ordinary	Drammensveien 288, Oslo, NO-0283, Norway
GlaxoSmithKline Australia Pty Ltd	Ordinary	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
GlaxoSmithKline B.V.	Ordinary	Van Asch van, Wijkstraat 55h, 3811 LP Amersfoort, The Netherlands, Netherlands
GlaxoSmithKline Beteiligungs GmbH	Ordinary	Prinzregentenplatz 9, 81675, Munchen, Germany
GlaxoSmithKline Biologicals Kft.	Ordinary	2100 Gödöllő, Homoki Nagy István utca 1, Hungary
GlaxoSmithKline Biologicals S.A.S.	Ordinary	637 Rue des Aulnois, Saint-Amand Les Eaux, 59230, France
GlaxoSmithKline Biologicals SA	Ordinary: Preference	Rue de l'Institut 89 B-1330 Rixensart, Belgium

Other statutory disclosures continued

Group companies continued

Name	Security	Registered address
Wholly owned subsidiaries continued		
GlaxoSmithKline Brasil Limitada	Quotas	Estrada dos Banderiantes, 8464, Rio de Janeiro, 22783-110, Brazil
GlaxoSmithKline Capital Inc.	Common	Wilmington Trust SP Services, Inc., 1100 N. Market Street, 4th Floor, Wilmington DE 19890, United States
GlaxoSmithKline Capital plc	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Caribbean Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Chile Farmaceutica Limitada	Social Capital	Av. Andrés Bello 2457, Torre 2, piso 20, Providencia, Santiago, Región Metropolitana, Chile
GlaxoSmithKline Colombia S.A.	Ordinary	Avenida El Dorado, #69B-45/Piso 9, Bogota, Colombia
GlaxoSmithKline Consumer Holding B.V. (ii)	Ordinary	Van Asch van Wijkstraat 55h, 3811 LP, Amersfoort, Netherlands
GlaxoSmithKline doo Beograd-Novi Beograd (in liquidation)	Ordinary	Milutin Milankovic, 1J, Novi Beograd, Belgrade, 11070, Serbia
GlaxoSmithKline Ecuador S.A.	Ordinary	Av 10 De Agosto N36-239, y Naciones Unidas, Edificio Electrocuatoriana, 2 do piso, Quito, Ecuador
GlaxoSmithKline El Salvador S.A. de C.V.	Ordinary	Municipio de San Salvador, Departamento de San Salvador, El Salvador
GlaxoSmithKline EOOD	Ordinary	16 Nedelcho Bonchev str., Sofia, Sofiya, 1592, Bulgaria
GlaxoSmithKline Export Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Export Panama S.A.	Ordinary	Panama City, Republic of Panama, Panama
GlaxoSmithKline Far East B.V.	Ordinary	Van Asch van Wijkstraat 55h, 3811 LP, Amersfoort, Netherlands
GlaxoSmithKline Finance plc	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline GmbH & Co. KG	Partnership Capital	Prinzregentenplatz 9, 81675, Munchen, Germany
GlaxoSmithKline Guatemala S.A.	Ordinary	3ra. Av. 13-78 Zona 10, Torre Citibank, Nivel 8, Guatemala City, Guatemala
GlaxoSmithKline Holding AS	Ordinary	Drammensveien 288, Oslo, NO-0283, Norway
GlaxoSmithKline Holdings (Americas) Inc.	Common	Wilmington Trust SP Services Inc., 1100 North Market Street, 4th Floor, Wilmington, Delaware, 19890
GlaxoSmithKline Holdings (One) Limited (i)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Holdings Limited (i)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Holdings Pty Ltd	Ordinary	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
GlaxoSmithKline Honduras S.A.	Ordinary	Tegucigalpa, MDC, Honduras
GlaxoSmithKline IHC Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Ilaclari Sanayi ve Ticaret A.S.	Nominative	Büyükdere Caddesi No. 173, 1.Levent Plaza B Blok, 1.Levent, Istanbul, 34394, Turkey
GlaxoSmithKline Inc.	Class A Common; Class C Preference	100 Milverton Drive, Suite 800, Mississauga ON L5R 4H1, Canada
GlaxoSmithKline Insurance Ltd.	Ordinary	c/o Trinity Corporate Services Ltd., Trinity Hall, 43 Cedar Avenue, Hamilton, Hamilton, HM12, Bermuda
GlaxoSmithKline Intellectual Property (No.2) Limited	Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
GlaxoSmithKline Intellectual Property Development Limited	Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
GlaxoSmithKline Intellectual Property Holdings Limited	A Ordinary; B Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
GlaxoSmithKline Intellectual Property Limited	Deferred; Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
GlaxoSmithKline Intellectual Property Management Limited	Ordinary	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
GlaxoSmithKline Investigación y Desarrollo, S.L.	Ordinary	Severo Ochoa 2 Parque Tecnológico de Madrid, Tres Cantos, 28760, Madrid Spain
GlaxoSmithKline Investments Pty Ltd	Ordinary	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
GlaxoSmithKline K.K.	Ordinary	1-8-1 Akasaka Minato-ku, Tokyo, Japan
GlaxoSmithKline Korea Limited	Ordinary	9F LS Yongsan Tower, 92 Hangang-daero, Yongsangu, Seoul, 04386, Korea, Republic of
GlaxoSmithKline Latin America, S.A.	Ordinary	Panama City, Republic of Panama, Panama
GlaxoSmithKline Limited	Ordinary	23/F., Tower 6, The Gateway, 9 Canton Road, Tsimshatsui, Kowloon, Hong Kong
GlaxoSmithKline Limited (ii)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline LLC	LLC Interests	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
GlaxoSmithKline Manufacturing SpA	Ordinary	Viale dell'Agricoltura 7, 37135, Verona, Italy
GlaxoSmithKline Maroc S.A.	Ordinary	42-44 Angle Bd, Rachidi et Abou Hamed El Glaza, Casablanca, Morocco
GlaxoSmithKline Medical and Healthcare Products Kft	Ordinary	1062 Budapest, Andrássy ut 113, Hungary
GlaxoSmithKline Mercury Limited (i)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England

Other statutory disclosures continued

Group companies continued

Name	Security	Registered address
Wholly owned subsidiaries continued		
GlaxoSmithKline Mexico S.A. de C.V.	Ordinary A; Ordinary B	Av. Real Mayorazgo 130 Piso 20, Colonia Xoco, Alcaldia Benito Juárez, Ciudad de Mexico, 03330, Mexico
GlaxoSmithKline NZ Limited	Ordinary	Level 2 E.2, Generator @GridAKL, 12 Madden Street, Wynyard Quarter, Auckland, 1010, New Zealand
GlaxoSmithKline Oy	Ordinary	Parkkalankatu 20 A, Helsinki, 00180, Finland
GlaxoSmithKline Peru S.A.	Ordinary	Av. Víctor Andrés Belaúnde N°147, Vía Principal N°133, Piso 7, Distrito de San Isidro, Lima, Peru
GlaxoSmithKline Pharma A/S	Ordinary	Vallensbæk Company House III, Delta Park 37, DK-2665, Valle, Denmark
GlaxoSmithKline Pharma GmbH	Ordinary	Wienerbergstraße 7, Wien, 1100, Austria, Austria
GlaxoSmithKline Pharmaceutical Kenya Limited	Ordinary	P.O Box 78392-00507, Likoni Road, Nairobi, Kenya
GlaxoSmithKline Pharmaceutical Nigeria Limited	Ordinary	1 Industrial Avenue, Ilupeju, Ikeja, Lagos, PM B 21218, Nigeria
GlaxoSmithKline Pharmaceutical Sdn Bhd	Ordinary	HZ.01, Horizon Penthouse, 1 Powerhouse, 1, Persiaran Bandar Utama, Bandar Utama, 47800 Petaling Jaya, Selangor, Malaysia
GlaxoSmithKline Pharmaceuticals (Pvt) Ltd	Ordinary	121 Galle Road, Kaldemulla, Moratuwa, Sri Lanka
GlaxoSmithKline Pharmaceuticals Costa Rica S.A	Ordinary	Autopista Florencia del Castillo, kilómetro siete, Oficentro TerraCampus, edificio uno, cuarto piso, San Diego, Cartago, 30302, Costa Rica
GlaxoSmithKline Pharmaceuticals SA	Ordinary	Avenue Fleming 20, 1300 Wavre, Belgium
GlaxoSmithKline Pharmaceuticals Ukraine LLC	Chartered Capital	Pavla Tychyn avenue, 1-V, Kiev, 02152, Ukraine
GlaxoSmithKline Philippines Inc	Ordinary	23rd Floor, The Finance Centre, 26th Street Corner 9th Avenue, Bonifacio Global City, Taguig City, 1634, Philippines
GlaxoSmithKline Pte Ltd	Ordinary	23 Rochester Park, 139234, Singapore
GlaxoSmithKline Puerto Rico, Inc.	Common	CORPORATION SERVICE COMPANY PUERTO RICO INC., c/o RVM Professional Services, LLC, A4 Reparto Mendoza, Humacao, 00791, Puerto Rico
GlaxoSmithKline Republica Dominicana S.A.	Ordinary	Blue Mall Tower, Floor 23 Ave., Winston Churchill 95, Santo Domingo, Dominican Republic
GlaxoSmithKline Research & Development Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline S.A.	Ordinary	Severo Ochoa, 2, Parque Tecnológico de Madrid, Tres Cantos, 28760, Madrid, Spain
GlaxoSmithKline S.p.A.	Ordinary	Viale dell'Agricoltura 7, 37135, Verona, Italy
GlaxoSmithKline s.r.o.	Ordinary	Hvezdova 1734/2c, Prague, 4 140 00, Czech Republic
GlaxoSmithKline Services GmbH & Co. KG	Partnership Capital	Prinzregentenplatz 9, 81675, Munchen, Germany
GlaxoSmithKline Services Unlimited (i)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Single Member A.E.B.E.	Ordinary	266 Kifissias Avenue, Halandri, Athens, 152 32, Greece
GlaxoSmithKline SL LLC	LLC Interests	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
GlaxoSmithKline SL LP (ii)(viii)	Partnership	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline South Africa (Pty) Limited	Ordinary	Flushing Meadows Building, The Campus, 57 Sloane Street, Bryanston 2021, South Africa
GlaxoSmithKline Trading Services Limited (iii)	Ordinary	12 Riverwalk, Citywest Business Campus, Dublin 24, Ireland
GlaxoSmithKline Tunisia S.A.R.L.	Ordinary	Immeuble REGUS, Lot B17, Centre Urbain Nord, Tunis, Tunisia
GlaxoSmithKline UK Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Uruguay S.A.	Registered Provisory Stock	Victor Soliño 349, Montevideo, Montevideo, 11300, Uruguay
GlaxoSmithKline US Trading Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GlaxoSmithKline Venezuela C.A.	Ordinary	calle Altagracia, edificio P&G, piso Mezzanina, torre Torre Sur, Urbanizacion Sorokaima, La Trinidad, Caracas, 1080, Venezuela, Bolivarian Republic of
GlaxoSmithKline Vietnam Limited Liability Company (ii)	Equity Capital	The Metropolitan, 235 Dong Khoi Street, District 1, 7th Floor Unit 701, Ho Chi Minh City, Vietnam
GlycoVaxyn AG (In liquidation)	Common; Preferred A; Preferred B; Preferred C	Grabenstrasse 3, 8952 Schlieren, Switzerland
Groupe GlaxoSmithKline	Ordinary	23 rue François Jacob, 92500, Rueil-Malmaison, France
GSK Biopharma Argentina S.A.	Nominative Non Endorseable Ordinary	Tucumán 1, piso 4, Buenos Aires, C1049AAA, Argentina
GSK (No.1) Scottish Limited Partnership (viii)	Partnership	50 Lothian Road, Festival Square, Edinburgh, Scotland, EH3 9WJ, United Kingdom
GSK (No.2) Scottish Limited Partnership (viii)	Partnership	50 Lothian Road, Festival Square, Edinburgh, Scotland, EH3 9WJ, United Kingdom
GSK (No.3) Scottish Limited Partnership (viii)	Partnership	50 Lothian Road, Festival Square, Edinburgh, Scotland, EH3 9WJ, United Kingdom

Other statutory disclosures continued

Group companies continued

Name	Security	Registered address
Wholly owned subsidiaries continued		
GSK Business Service Centre Sdn Bhd	Ordinary	Level 6, Quill 9, 112 Jalan Prof. Khoo Kay Kim, Petaling Jaya, 46300 Selangor, Malaysia
GSK Capital B.V. (iii)(v)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS
GSK Capital K.K.	Ordinary	1-8-1 Akasaka Minato-ku, Tokyo, Japan
GSK Commercial Sp. z o.o.	Ordinary	ul. Rzymowskiego 53, 02-697, Warsaw, Poland
GSK d.o.o., Ljubljana	Ordinary	Ameriška ulica 8, Ljubljana, 1000, Slovenia
GSK Enterprise Management Co, Ltd	Ordinary	Floor 4, 18 Lane 999 Huanke Road, No. 1358 Zhongke Road, Shanghai, China
GSK Equity Investments, Limited	Units	Corporation Service Company, 2595 Interstate Drive, Suite 103, Harrisburg PA 17110, United States
GSK Finance (No.3) PLC	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GSK Finance (No 2) Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
GSK India Global Services Private Limited	Equity	Level 1, 2 & 3 Luxor North Tower, Bagmane Capital Business Park Outer Ring Road, Bangalore, Karnataka, 560037, India
GSK International Holding and Finance BV	Ordinary	Van Asch van Wijckstraat 55h, 3811 LP, Amersfoort, Netherlands
GSK Kazakhstan LLP	Participation Interest	Nursultan Nazarbayev Ave 273, Business center USKO, 3rd fl., Almaty, 050059, Kazakhstan
GSK Life Sciences FZE	Ordinary	LB06015, Jebel Ali Freezone, Dubai, United Arab Emirates
GSK Pharma India Private Limited	Equity	1, Battery House, Bhulabhai Desai Road, Mumbai, Maharashtra, 400026, India
GSK Pharma Vietnam Company Limited	Chartered Capital	Unit 702/703 7th Floor, The Metropolitan Tower, 235 Dong Khoi Street, Ben Nghe Ward, District 1, Ho Chi Minh, Vietnam
GSK Pharmaceutical Trading S.A. (ii)	Ordinary	Bucharest, 1-5 Costache Negri Street, Opera Center One, 5th floor, discussions room 01, District 5, Romania
GSK PSC Poland sp. z o.o.	Equal and indivisible shares	ul. Grunwaldzka 189, Poznań, 60-322, Pol
GSK Services Sp z o.o.	Ordinary	Ul. Grunwaldzka 189, 60-322, Poznan, Poland
GSK Vaccines BV	Ordinary	Hullenbergweg 85, 1101 CL, Amsterdam, Netherlands
GSK Vaccines GmbH	Ordinary	Emil-von-Behring-Str.76, 35041 Marburg, Germany
GSK Vaccines Institute for Global Health S.r.l.	Quota	Via Fiorentina 1, 53100, Siena, Italy
GSK Vaccines S.r.l.	Quota	Via Fiorentina 1, 53100, Siena, Italy
GSK Vaccines Vertriebs GmbH	Ordinary	Rudolf-Diesel-Ring 27, 83607, Holzkirchen, Germany
Human Genome Sciences, Inc.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
ID Biomedical Corporation of Quebec	Common	2323, boul. Du Parc Technologique, Québec Québec GIP 4R8, Canada
Instituto Luso Farmaco, Limitada (ii)	Quota	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
InterPharma Dienstleistungen GmbH	Quota	Wienerbergstraße 7, Wien, 1100, Austria, Austria
J&J Technologies, LC (ii)	LLC Interests	Corporation Service Company, 100 Shockoe Slip, 2nd Floor, Richmond VA 23219, United States
JSC GlaxoSmithKline Trading	Ordinary	Leningradskiy Prospect 37A, Building 4, Floor 3, Premises XV, Room 1, 125167, Moscow, Russian Federation
Laboratoire GlaxoSmithKline	Ordinary	23 rue François Jacob, 92500, Rueil-Malmaison, France
Laboratoire Pharmaceutique Algérien LPA Production SPA	Ordinary	Zone Industrielle Est, Boudouaou, Boumerdes, Algeria
Laboratoire Pharmaceutique Algérien SPA	Ordinary	Zone Industrielle Est, Boudouaou, Boumerdes, Algeria
Laboratoires Paucourt (ii)	Ordinary	23 rue François Jacob, 92500, Rueil-Malmaison, France
Laboratoires Saint-Germain (ii)	Ordinary	23 rue François Jacob, 92500, Rueil-Malmaison, France
Laboratorios Dermatologicos Darier, S.A de C.V.	Ordinary A; Ordinary B	Av. Real Mayorazgo 130 Piso 20, Colonia Xoco, Alcaldia Benito Juárez, Ciudad de Mexico, 03330, Mexico
Laboratorios Farmaceuticos Stiefel (Portugal) LTDA (ii)	Ordinary	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
Laboratorios Stiefel de Venezuela SA	Ordinary	Calle Altagracia, edificio P&G, nivel Mezzanina., piso Mezzanina, local Torre Sur, Urbanizacion Sorokaima, La Trinidad, Caracas, 1080, Venezuela, Bolivarian Republic of
Laboratorios Stiefel Ltda.	Ordinary	Rua Professor Joao Cavalheiro Salem, no.1077, Bairro de Bonsucesso, Municipality of Guarulhos, Sao Paulo, CEP 07243-580, Brazil
Laboratorios Wellcome De Portugal Limitada (ii)	Quota	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
Maxinutrition Limited (in liquidation)	Ordinary	C/O BDO LLP, 5 Temple Square, Temple Street, Liverpool, L2 5RH
PT Glaxo Wellcome Indonesia	Class A; Class B	JL. Pulobuaran Raya Kav.III/ DD 2,3,4 KWS. Industri, Pulogadung, Jatinegara, Cakung, Jakarta Timur, Indonesia
Setfirst Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England

Other statutory disclosures continued

Group companies continued

Name	Security	Registered address
Wholly owned subsidiaries continued		
Sierra Oncology Australia Pty Ltd	Ordinary	c/o Maddocks Lawyers, Angel Place, Level 27, 123 Pitt Street Sydney 2000, Australia
Sierra Oncology Canada ULC	Common	Suite 1800 - 510 West Georgia Street, Vancouver BC V6B 0M3, Canada
Sitari Pharma, Inc.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Smith Kline & French Portuguesa-Produtos Farmaceuticos, LDA (ii)	Ordinary	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflores, 1495-131, Alges, Portugal
SmithKline Beecham (Bangladesh) Private Limited (ii)	Ordinary	House-2/A, Road-138, Gulshan-1, Dhaka, 1212, Bangladesh
SmithKline Beecham (Cork) Limited	Ordinary	12 Riverwalk, Citywest Business Campus, Dublin 24, Ireland
SmithKline Beecham Egypt L.L.C.	Quota	Amoun Street, El Salam City, Cairo, Egypt
SmithKline Beecham Farma, S.A.	Ordinary	Severo Ochoa, 2, Parque Tecnologico de Madrid, Tres Cantos, 28760, Madrid, Spain
SmithKline Beecham Legacy H Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
SmithKline Beecham Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
SmithKline Beecham Pension Plan Trustee Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
SmithKline Beecham Pharma GmbH & Co KG	Partnership Capital	Prinzregentenplatz 9, 81675, Munchen, Germany
SmithKline Beecham Pharma Verwaltungs GmbH	Ordinary	Prinzregentenplatz 9, 81675, Munchen, Germany
SmithKline Beecham Pharmaceuticals (Pty) Limited (ii)	Ordinary	Flushing Meadows Building, The Campus, 57 Sloane Street, Bryanston 2021, South Africa
SmithKline Beecham Pharmaceuticals Co.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
SmithKline Beecham Senior Executive Pension Plan Trustee Limited (ii)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Stiefel GmbH & Co. KG	Partnership Capital	Prinzregentenplatz 9, 81675, Munchen, Germany
Stiefel Laboratories Legacy (Ireland) Limited	Ordinary	Unit 2 Building 2500, Avenue 2000 Cork Airport Business Park, Cork, Ireland
Stiefel Laboratories Pte Limited	Ordinary	1 Pioneer Sector, 628413, Singapore
Stiefel Laboratories, Inc.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Stiefel Maroc SARL	Ordinary	275 Boulevard Zerktoni, Casablanca, Morocco
Stiefel Research (Australia) Holdings Pty Ltd	Ordinary	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
Stiefel Research Australia Pty Ltd	Ordinary	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
Stiefel West Coast LLC	LLC Interests	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Strebor Inc.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Tesaro Bio GmbH (in liquidation)	Ordinary	Poststrasse 6, 6300 Zug, Switzerland
Tesaro Bio Netherlands B.V	Ordinary	Joop Geesinkweg 901, 1114 AB, Amsterdam-Duivendrecht, Netherlands
Tesaro Development, Ltd.	Ordinary	Clarendon House, 2 Church Street, Hamilton HM11, Bermuda
Tesaro, Inc.	Common	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
The Sydney Ross Co. (ii)	Ordinary	Corporation Service Company, Princeton South Corporate Center, Suite 160, 100 Charles Ewing Blvd, Ewing NJ 08628, United States
UCB Pharma Asia Pacific Sdn Bhd (ii)	Ordinary	12th Floor, Menara Symphony, No. 5, Jalan Prof. Khoo Kay Kim, Seksyen 13, 46200 Petaling Jaya, Malaysia
Wellcome Consumer Healthcare Limited (ii)	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Wellcome Limited	Ordinary	980 Great West Road, Brentford, Middlesex, TW8 9GS, England

Name	Security	Effective % Ownership	Registered address
Subsidiaries where the effective interest is less than 100%			
Amoun Pharmaceutical Industries Co. S.A.E.	Monetary Shares	90.71%	El Salam City 11491, PO Box 3001, Cairo, Egypt
Biddle Sawyer Limited	Equity	75.00%	252 Dr Annie Besant Road, Mumbai, 400030, India
British Pharma Group Limited (i)	Guarantee (50%)	50.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
Galvani Bioelectronics Inc.	Common	55.00%	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
Galvani Bioelectronics Limited	A Ordinary; B Ordinary	55.00% -	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
Glaxo Laboratories (Nigeria) Limited (ii)	Ordinary	99.99%	82 Marine Road, Apapa, Lagos, Nigeria
Glaxo-Allenburys (Nigeria) Limited (ii)	Ordinary	99.00%	41 Creek Road, Apapa, Lagos, PMB 1401, Nigeria

Other statutory disclosures continued

Group companies continued

Name	Security	Effective % Ownership	Registered address
Subsidiaries where the effective interest is less than 100% continued			
GlaxoSmithKline (Tianjin) Co. Ltd	Ordinary	90.00%	No. 65, the Fifth Avenue, Tai Feng Industrial Park, Tianjin Economic and Technolog, Tianjin, 300457, China
GlaxoSmithKline Algérie S.P.A.	Ordinary	99.99%	Zone Industrielle Est, Boudouaou, Wilaya de Boumerdes, Algeria
GlaxoSmithKline Consumer Nigeria plc (vi)	Ordinary	46.42%	1 Industrial Avenue, Ilupeju, Ikeja, Lagos, PM B 21218, Nigeria
GlaxoSmithKline Pakistan Limited	Ordinary	82.59%	The Sykes Building, 35 Dockyard Road, West Wharf, Karachi, 74000, Pakistan
GlaxoSmithKline Pharmaceuticals Limited	Equity	75.00%	252 Dr Annie Besant Road, Mumbai, 400030, India
GlaxoSmithKline S.A.E.	Ordinary	91.20%	Boomerang Office Building - Land No. 46, Zone (J) - 1st District, Town Center - 5th Tagammoe, New Cairo City, Egypt
Laboratorios ViiV Healthcare, S.L.	Ordinary	78.30%	Severo Ochoa, 2, Parque Tecnológico de Madrid, Tres Cantos, 28760, Madrid, Spain
Modern Pharma Trading Company L.L.C.	Quota	98.24%	Amoun Street, PO Box 3001, El Salam City, Cairo, 11491, Egypt
PHIVCO-1 LLC	LLC Interests	78.30%	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
PHIVCO-2 LLC	LLC Interests	78.30%	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
SmithKline Beecham-Biomed O.O.O.	Participation Interest	97.00%	Leningradskiy Prospect 37A, Building 4, Floor 2, Premises XIV, Room 42, 1251 67, Moscow, Russian Federation
Stiefel Egypt LLC (ii)	Quota	99.00%	Amoun Street, PO Box 3001, El Salam City, Cairo, 11491, Egypt
ViiV Healthcare (South Africa) (Proprietary) Limited	Ordinary	78.30%	Flushing Meadows Building, The Campus, 57 Sloane Street, Bryanston 2021, South Africa
ViiV HealthCare BV	Ordinary	78.30%	Van Asch van, Wijkstraat 55h, 3811 LP Amersfoort, The Netherlands, Netherlands
ViiV Healthcare Company	Common	78.30%	Corporation Service Company, 251 Little Falls Drive, Wilmington DE 19808, United States
ViiV Healthcare Finance 2 Limited	Ordinary	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
ViiV Healthcare Finance Limited	Ordinary; Redeemable Preference	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
ViiV Healthcare GmbH	Ordinary	78.30%	Prinzregentenplatz 9, 81675, Munchen, Germany
ViiV Healthcare GmbH	Ordinary	78.30%	Talstrasse 3, 3053 Muenchenbuchsee, Switzerland
ViiV Healthcare K.K.	Ordinary	78.30%	1-8-1 Akasaka Minato-ku, Tokyo, Japan
ViiV Healthcare Limited	A Ordinary; B Ordinary; C Ordinary; D1 Preference; D2 Ordinary; Deferred; E 5% Cumulative Preference	78.30%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
ViiV Healthcare Pty Ltd	Ordinary	78.30%	Level 4, 436 Johnston Street, Abbotsford, Victoria, 3067, Australia
ViiV Healthcare Puerto Rico, LLC	LLC Interests	78.30%	CORPORATION SERVICE COMPANY PUERTO RICO INC., c/o RVM Professional Services, LLC, A4 Reparto Mendoza, Humacao, Puerto Rico, 00791
ViiV Healthcare S.r.l.	Quota	78.30%	Viale dell'Agricoltura 7, 37135, Verona, Italy
ViiV Healthcare SAS	Ordinary	78.30%	23 rue François Jacob, 92500, Rueil-Malmaison, France
ViiV Healthcare sprl	Ordinary	78.30%	Avenue Fleming 20, 1300 Wavre, Belgium
ViiV Healthcare Trading LLC (ii)	Participation Interest	78.30%	Leningradskiy Prospect 37A, Building 4, Floor 2, Premises XIV, Room 28, 1251 67, Moscow, Russian Federation
ViiV Healthcare Trading Services UK Limited	Ordinary	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
ViiV Healthcare UK (No.3) Limited	Ordinary	78.30%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
ViiV Healthcare UK (No.4) Limited	Ordinary	78.30%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
ViiV Healthcare UK (No.5) Limited	Ordinary	78.30%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
ViiV Healthcare UK (No.6) Limited	Ordinary	78.30%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
ViiV Healthcare UK (No.7) Limited	Ordinary	78.30%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage, SG1 2NY, United Kingdom
ViiV Healthcare UK Limited	Ordinary	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England
ViiV Healthcare ULC	Common	78.30%	3500 855-2nd Street SW, Calgary AB T2P 4J8, Canada

Other statutory disclosures continued

Group companies continued

Name	Security	Effective % Ownership	Registered address
Subsidiaries where the effective interest is less than 100% continued			
ViiVHIV Healthcare Unipessoal Lda	Quota	78.30%	Rua Dr Antonio Loureiro Borges No 3, Arquiparque, Miraflares, 1495-131, Alges, Portugal
Winster Pharmaceuticals Limited	Ordinary	46.42%	2A Association Avenue, Ilupeju Industrial Estate, Lagos, PO Box 3199, Nigeria

Name	Security	Effective % Ownership	Registered address
Associates			
GlaxoSmithKline Landholding Company, Inc	Common	39.93%	23rd Floor, The Finance Centre, 26th Street Corner 9th Avenue, Bonifacio Global City, Taguig City, 1634, Philippines
Index Ventures Life VI (Jersey) LP	Partnership Interest (24.94%)	24.94%	44 Esplanade, St Helier, Jersey, JE4 9WG, Channel Islands
Kurma Biofund II FCPR	Partnership Interest (32.06%)	32.06%	24 rue Royale, 5th Floor, 75008, Paris, France
Longwood Fund I, LP	Partnership Interest (35%)	35.00%	The Prudential Tower, Suite 1555, 800 Boylston Street, Boston, MA 02199
Medicxi Ventures I LP	Partnership Interest (26.10%)	26.10%	44 Esplanade, St Helier, Jersey, JE4 9WG, Channel Islands

Joint Ventures

Chiron Panacea Vaccines Private Limited (in Liquidation)	Equity Shares	50.00%	708/718, 7th Floor, A Wing, Sagar Tech Plaza, Saki Naka, Andheri East, Mumbai, Maharashtra, 400072, India
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Other significant holdings

Axon Therapies, Inc	Common (2.63%); Series A Preference (18.40%)	20.03%	2326 Walsh Avenue Santa Clara, CA 95051, United States
Alpheus Medical, Inc.	Series A Preference (13.77%); Series A-1 Preference (7.27%)	21.04%	3510 Hopkins Place, North Oakdale, Minnesota 55128, USA
Global Farm S.A.	A Shares (0%); B Shares (0%); C Shares (100%)	20.00% 100% of C Shares	Mendoza 1259, Ciudad Autónoma de Buenos Aires, Argentina
Longwood Fund II, LP	Partnership Interest (20.00%)	20.00%	The Prudential Tower, Suite 1555, 800 Boylston Street, Boston, MA 02199
Sanderling Ventures VII, L.P. A63	Partnership Interest (25.31%)	25.31%	400 S. El Camino Real, Suite 1200, San Mateo, CA 94402
SR One Capital Fund I-B, LP	Partnership Interest (44%)	44.00%	Corporation service company, 251 Little Falls Drive, City of Wilmington, County of New Castle, Delaware 19808
SR One Capital Opportunities Fund I, LP	Partnership Interest (24.46%)	24.46%	Corporation service company, 251 Little Falls Drive, City of Wilmington, County of New Castle, Delaware 19808

Other statutory disclosures continued

Group companies continued

The following UK registered subsidiaries will take advantage of the audit exemption set out within Section 479A of the Companies Act 2006 for the period ended 31 December 2023. Unless otherwise stated, the undertakings listed below are owned, either directly or indirectly, by GSK plc.

Name	Security	Effective % Ownership	Registered address	Company Number
UK registered subsidiaries exempted from audit				
Burroughs Wellcome International Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	543757
Domantis Limited	Ordinary	100.00%	GSK Medicines Research Centre, Gunnels Wood Road, Stevenage SG1 2NY, United Kingdom	3907643
Edinburgh Pharmaceutical Industries Limited (ii)	Ordinary; Preference;	100.00%	Shewalton Road, Irvine, Ayrshire, KA11 5AP, United Kingdom	SC005534
Eskaylab Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	99025
Glaxo Wellcome UK Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	480080
Glaxochem (UK) Unlimited	Ordinary; Ordinary B; Ordinary C	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	4299472
GlaxoSmithKline Intellectual Property (No.3) Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	11480952
GlaxoSmithKline Intellectual Property (No.4) Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	11721880
GlaxoSmithKline Intellectual Property (No.5) Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	11959399
GlaxoSmithKline International Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	2298366
GSK GP 1 Limited (iv)	A Shares; B Shares	100.00%	50 Lothian Road, Festival Square, Edinburgh, Scotland, EH3 9WJ, United Kingdom	SC721605
GSK GP 2 Limited (iv)	Ordinary	100.00%	50 Lothian Road, Festival Square, Edinburgh, Scotland, EH3 9WJ, United Kingdom	SC721606
GSK LP Limited (i)(iv)	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	13879411
Montrose Fine Chemical Company Ltd.	Ordinary	100.00%	Shewalton Road, Irvine, Ayrshire, KA11 5AP, United Kingdom	SC190635
PHIVCO UK II Limited	Ordinary	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	6944229
PHIVCO UK Limited	Ordinary	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	6944223
Smith Kline & French Laboratories Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	52207
SmithKline Beecham (Export) Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	2860752
SmithKline Beecham (H) Limited	Non-cumulative Non-redeemable; Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	3296131
SmithKline Beecham (Investments) Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	302065
SmithKline Beecham Marketing and Technical Services Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	494385
SmithKline Beecham Nominees Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	503868
SmithKline Beecham Overseas Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	2552828
SmithKline Beecham Pension Plan Trustee Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	3425311
Stiefel Laboratories (U.K.) Ltd	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	831160
Tesaro UK Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	7890847
The Wellcome Foundation Limited	Ordinary	100.00%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	194814
ViiV Healthcare Overseas Limited	Ordinary	78.30%	980 Great West Road, Brentford, Middlesex, TW8 9GS, England	7027385

In accordance with Section 479C of the Companies Act 2006, the company will guarantee debts and liabilities of the above UK subsidiary undertakings. As at 31 December 2023 the total sum of these debts and liabilities is £317 million (2022 – £1266 million)

Key

- (i) Directly owned by GSK plc.
- (ii) Dormant entity.
- (iii) Tax resident in the UK.
- (iv) Exempt under Regulation 7 of the Partnership (Accounts) Regulations 2008 from the requirement to deliver to the registrar financial statements of the qualifying partnership(s) of which the entity is a member in accordance with the Companies Act.
- (v) Incorporated in the Netherlands
- (vi) Consolidated as a subsidiary in accordance with Section 1162 (4)(a) of the Companies Act 2006 on the grounds of dominant influence.
- (vii) Principal business address in Puerto Rico.
- (viii) Exempt from the provisions of Regulations 4-6 of the Partnership (Accounts) Regulation 2008, in accordance with the exemptions noted in Regulation 7 of that Regulation.

Glossary of terms

Terms used in the Annual Report	US equivalent or brief description
Accelerated capital allowances	Tax allowance in excess of depreciation arising from the purchase of fixed assets that delay the charging and payment of tax. The equivalent of tax depreciation.
American Depositary Receipt (ADR)	Receipt evidencing title to an ADS. Each GSK ADR represents two Ordinary Shares
American Depositary Shares (ADS)	Listed on the New York Stock Exchange; represents two Ordinary Shares
Basic earnings per share	Basic income per share
Called up share capital	Ordinary Shares, issued and fully paid.
CER growth	Growth at constant exchange rates.
The company	GSK plc
Currency swap	An exchange of two currencies, coupled with a subsequent re-exchange of those currencies, at agreed exchange rates and dates
Defined benefit plan	Pension plan with specific employee benefits, often called 'final salary scheme'.
Defined contribution plan	Pension plan with specific contributions and a level of pension dependent upon the growth of the pension fund.
Derivative financial instrument	A financial instrument that derives its value from the price or rate of some underlying item
Diluted earnings per share	Diluted income per share.
Employee Share Ownership Plan Trusts	Trusts established by the Group to satisfy share-based employee incentive plans
Equity Shareholders' funds	Shareholders' equity.
Finance lease	Capital lease.
Freehold	Ownership with absolute rights in perpetuity
The Group	GSK plc and its subsidiary undertakings.
GSK	GSK plc and its subsidiary undertakings.
Hedging	The reduction of risk, normally in relation to foreign currency or interest rate movements, by making off-setting commitments.
Intangible fixed assets	Assets without physical substance, such as computer software, brands, licences, patents, know-how and marketing rights purchased from outside parties.
Ordinary share	A fully paid up ordinary share in the capital of the company.
Profit	Income
Profit attributable to shareholders	Net income
Share capital	Ordinary Shares, capital stock or common stock issued and fully paid.
Share option	Stock option.
Share premium account	Additional paid-up capital or paid-in surplus (not distributable).
Shares in issue	The number of shares outstanding.
Subsidiary	An entity in which GSK exercises control.
Treasury share	Treasury stock.
Turnover	Revenue.
UK Corporate Governance Code	As required by the UK Listing Authority, the company has disclosed in the Annual Report how it has applied the best practice corporate governance provisions of the Financial Reporting Council's UK Corporate Governance Code.