## SASB INDEX 2021



## SASB Standard: Biotechnology & Pharmaceuticals

This report is an index to the location of our disclosures that align with the Sustainability Accounting Standards Board (SASB) standards for Biotechnology & Pharmaceuticals. The report provides data from 1 January 2021 to 31 December 2021, unless otherwise stated.

Metric Code	Metric	Disclosure Location
Safety of Clinical	Trial Participants	
HC-BP-210a.1	Discussion, by world region, of management process for ensuring quality and patient safety during clinical trials	<u>2021 Annual Report</u> , p. 34 <u>Global Standard: Bioethics</u> , pp.4-7
HC-BP-210a.2	Number of FDA Sponsor Inspections related to clinical trial management and pharmacovigilance that resulted in: (1) Volun- tary Action Indicated (VAI) and (2)Official Action Indicated (OAI)	2021: VAI - 0, OAI - 0 2020: VAI - 0, OAI - 0 2019: VAI - 0, OAI - 0
HC-BP-210a.3	Total amount of monetary losses as a result of legal proceedings associated with clinical trials in developing countries	Not reported
Access to Medici	nes	
HC-BP-240a.	Description of actions and initiatives to promote access to health care products for priority diseases and in priority countries as defined by the Access to Medicine Index	Access to Medicine Index 2021 Report, pp. 136-139
HC-BP-240a.2	List of products on the WHO List of Prequalified Medicinal Products as part of its Prequalification of Medicines Programme (PQP)	Not reported
Affordability and	Pricing	
HC-BP-240b.1	Number of settlements of Abbreviated New Drug Application (ANDA) litigation that involved payments and/or provisions to delay bringing an authorized generic product to market for a defined time period	Not reported
HC-BP-240b.2	Percentage change in: (1) average list price and (2) average net price across U.S. product portfolio compared to previous year	Not reported
HC-BP-240b.3	Percentage change in: (1) list price and (2) net price of product with largest increase compared to previous year	Not reported
Drug Safety		
HC-BP-250a.1	List of products listed in the Food and Drug Administration's (FDA) MedWatch Safety Alerts for Human Medical Products database	FDA Adverse Event Reporting webpage
HC-BP-250a.2	Number of fatalities associated with products as reported in the FDA Adverse Event Reporting System	FDA Adverse Event Reporting webpage
HC-BP-250a.3	Number of recalls issued, total units recalled	2021 Sustainability Data Summary, p. 15
HC-BP-250a.4	Total amount of product accepted for take-back, reuse, or disposal	Not reported
HC-BP-250a.5	Number of FDA enforcement actions taken in response to violations of current Good Manufacturing Practices (cGMP), by type	FDA - Inspection Citations; FDA- Warning <u>letters</u>
Counterfeit Drugs	s	
HC-BP-260a.1	Description of methods and technologies used to maintain traceability of products throughout the supply chain and prevent counterfeiting	SASB response: Counterfeit drugs
HC-BP-260a.2	Discussion of process for alerting customers and business partners of potential or known risks associated with counterfeit products	SASB response: Counterfeit drugs
HC-BP-260a.3	Number of actions that led to raids, seizure, arrests, and/or filing of criminal charges related to counterfeit products	Not Reported

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Metric Code	Metric	Disclosure Location		
Ethical Marketing	J			
HC-BP-270a.1	Total amount of monetary losses as a result of legal proceedings associated with false marketing claims	Not Reported		
HC-BP-270a.2	Description of code of ethics governing promotion of off-label use of products	AstraZeneca Global Standard: Promoting our products		
Employee Recruitment, Development and Retention				
HC-BP-330a.1	Discussion of talent recruitment and retention efforts for scientists and research and development personnel	2021 Annual Report, pp. 41-42		
HC-BP-330a.2	(1)Voluntary and (2) involuntary turnover rate for: (a) executives/senior managers, (b) mid-level managers, (c) professionals, and (d) all others	2021 Sustainability Data Summary, p. 15		
Supply Chain Management				
HC-BP-430a.1	Percentage of (1) entity's facilities and (2) Tier I suppliers' facilities participating in the Rx-360 International Pharmaceutical Supply Chain Consortium audit program or equivalent third-party audit programs for integrity of supply chain and ingredi- ents	We do not use third-party auditing organisations, AstraZeneca has an internal risk based supplier assessment system. In 2021 Global Quality Audit completed 425 total audits consisting of 379 supplier au- dits, 29 AstraZeneca internal audits and 17 Marketing Company audits		
Business Ethics				
HC-BP-510a.1	Total amount of monetary losses as a result of legal proceedings associated with corruption and bribery	Not Reported		
HC-BP-510a.2	Description of code of ethics governing interactions with health care professionals	AstraZeneca and Global Transparency		
Activity Metrics				
HC-BP-000.A	Number of patients treated	Not Reported		
HC-BP-000.B	Number of drugs (1) in portfolio and (2) in research and development (Phases 1-3)	2021 Annual Report		