IS4H TOOLKIT
KNOWLEDGE CAPSULES
DATA DISAGGREGATION IN PUBLIC HEALTH

DEPARTMENT OF EVIDENCE AND INTELLIGENCE FOR ACTION IN HEALTH
PAHO/WHO
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Information Systems for Health Toolkit

Knowledge Capsules
Understanding Data Disaggregation in Public Health

IS4H-KCDD

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Department of Evidence and Intelligence for Action in Health
Pan American Health Organization - World Health Organization
Data disaggregation in Public Health

The monitoring, evaluation and analysis of the health status of the population is one of the essential public health functions of national health authorities. This includes the monitoring of the population health and trends including its determinants and identifying inequities in risks, threats and access to services based on data that national information systems are routinely collecting. In this regard, the availability of data by various dimensions (disaggregation) is of utmost importance to be able to capture health inequalities that exist within a population.

The 2030 Agenda for Sustainable Development that was approved in 2015 and includes an overarching principles of data disaggregation, face some of the challenges related to the data since in many occasions, not all the population is counted due to the deficiencies in the information systems of the countries, which hinders progress. For this reason, having solid and accurate data is essential for targeting initiatives, which means that data disaggregation plays an important role in this process.

What does data disaggregation mean?

Disaggregated data refers to the separation of compiled information into smaller units to elucidate underlying trends and patterns. Compiled data may come from multiple sources (the public/private sectors and national/international organizations) and have multiple variables or “dimensions.” To enhance understanding of a situation, the data is grouped by dimension, such as age, sex, geographic area, education, ethnicity, or other socioeconomic variables.

What are some of the benefits of data disaggregation?

In 2015, PAHO Member States recognized that the basic principle of the 2030 Agenda — that no one is left behind — will require a significant level of data disaggregation. In alignment with the SDGs, many countries are working to strengthen their national capacity to improve the collection of disaggregated health information to better inform policy, strategy, and operational and clinical decision-making through the identification of factors that make some populations more vulnerable. High quality, accessible, trusted, timely, open and reliable disaggregated data can lead to many benefits, such as strengthened Health Information Systems that allow:

- Accurate health situation analysis
- Enhanced understanding of a population’s particular characteristics
- Problem detection
- Patterns and needs identification
- Equity monitoring
- Well-founded plans, policies and strategies
- Real-time decision making
- Financing structures and plans for resource targeting
- Project monitoring and evaluation
- Progress measurement
- Routine comparisons and trends analyses to inform and improve programs

**Interoperability is key to analyze disaggregated data across national databases, registries and information systems** *(see related capsule)*

**What is the minimum set of suggested disaggregation?**

Once data is collected, there are many ways to divide it in different units, depending on the information available and the objectives sought. In accordance with the Fundamental Principles of Official Statistics, indicators should be disaggregated by:5,6

![Hexagonal diagram](hexagonal_diagram.png)
**What are the principles of data disaggregation?**

The Global Partnership for Sustainable Development Data sets for the following criteria:

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**Principles of Inclusive Data Charter**

1. All populations must be included in the data.
2. All data should, wherever possible, be disaggregated in order to accurately describe all populations.
3. Data should be drawn from all available sources.
4. Those responsible for the collection of data and production of statistics must be accountable.
5. Human and technical capacity to collect, analyze, and use of disaggregated data must be improved, including through adequate and sustainable financing.


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**What are the mandates associated with data disaggregation?**

Some of the mandates compromised with data disaggregation include:

- The 2030 Agenda for Sustainable Development
- Information Systems for Health (IS4H) Strategic Plan
- Vital Statistics Plan of Action
- The Sustainable Health Agenda for the Americas 2018-2030
Where can I find technical information about data disaggregation?

4. UNSD. Background document. Data disaggregation and SDG Indicators. IAEG-SDGs, 2019.