Despite these problems, South Africa has managed to develop national standards that are flexible enough to "absorb" local innovations and changes over time. The following points may help to explain why this is so.

- The “hierarchy of standards” ("information needs" in Shaw’s figure) has been a powerful tool to negotiate a balance between the national needs for control with the local (e.g. province or health programme) needs for flexibility or more data. While all health units are required to collect and report the core national data, they are at the same time allowed to collect their own additional data.
- Use of information is highlighted by linking data sets to targets and indicators.
- The flexible approach to standards following the hierarchy makes it easy to absorb and implement changes over time; there is no “final” data set. Local innovations are allowed for and may eventually be included in the national data set.
- The flexibility of the South African District Health Information System (DHIS) database application is crucial to managing the ever changing national and local data sets. Data elements, indicators and data sets are added, edited and managed by the health services themselves, thus making it possible to manage multiple data sets at district level. This “data repository” or “warehouse” approach may be a key to how the lessons from South Africa could be applied in other countries.

Over the years, I have been involved in efforts to apply the South African lessons in many countries. It has not been easy. National health information system databases and reporting formats tend to be rigid and unable to respond to changes, thus leading to fragmentation of the system. The HIV/AIDS programmes are currently aggravating this situation.

So what can be done? Current efforts to establish integrated data sets in contexts as different as Addis Ababa (Ethiopia), Botswana, Zanzibar (United Republic of Tanzania), Andhra Pradesh (India) and Ho Chi Minh City (Viet Nam) may provide some answers. Here data sets from all or most programmes are combined and streamlined by sorting out overlaps, gaps and inconsistencies. Following the South African district data warehouse approach, the combined essential data set is then further improved and reduced by focusing on the need-to-know indicators. Programme-specific software applications are linked electronically to the DHIS, thus providing a shared data repository. The objective of integrating all indicators relevant to the Millennium Development Goals has proven important in building consensus.

Purists may argue that the data warehouse approach based on flexible standards advocated here is violating the spirit of the South African minimum essential data set approach, by taking a combined — maximum — data set as the point of departure. It may, however, be the most appropriate way to apply the South African indicator-driven approach in a situation increasingly dominated by strong programmes and multiple uncoordinated data sets and software applications.

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The paper by Vincent Shaw highlights some of the long awaited practical approaches to ensure relevance and use of health information systems in developing countries. The South African experience he recounts is very relevant as current “best practice”. It is a very important step forward in the simplification and integration of programme areas and routine services into a shared essential set of routine data. The cohesion of health services and the whole system will be much strengthened by such a shared data reference and information base.

The essential data are still to be selected by programme managers, however, even though the definition of the data set emanated from the district level. It is not shown how the data will be turned into useful information that will assist in planning and monitoring at all levels of the health system. The examples taken from the nutrition programme and the Expanded Programme on Immunization only exemplify health status (outcome) and service provision (output) data.

In the section on specific programme surveys, resource and staff availability are mentioned as well as service quality. It is worrying that these are not shown to be included in an essential routine data set, as it is extremely important that an essential data set allows production and sharing of information on health management. It is also not shown whether quality of care will be viewed from both provider and user perspectives and whether user views on service priorities and other qualitative data are included.

In Shaw’s figure, the column of information used by the national level on a routine basis within the triangle of information should be seen as the core data on facility performance that must be shared in an accessible database for the whole health service and other parts of the health system, and be available for sharing with users and the public.

The information needs triangle could also be depicted as service quality at the bottom, supervision and coordination needs at the intermediate level, and policy needs at the top. The main focus of the data set must be on service quality, with less emphasis on supervision and coordination and even less on policy. Service quality must relate to health management, service output and outcomes as viewed from both the provider and the user sides.

The number of data elements and indicators still seems to be high in relation to similar elements in the core part of the health information system in other African countries and may indicate a still limited degree of compromise and shared focus between the levels and programmes. A stronger emphasis is needed on the iterative nature of the health information system, and not so much on annual or quarterly “reporting”. We need to move away from a culture of reporting to a culture of using the data for ours flights or districts first — on a continuous basis. Benchmarking and quality assurance processes exemplify some useful frameworks for continuous use of data.

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