A systematic review of counterfeit and substandard medicines in field quality surveys

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Objectives: Counterfeit and substandard medicines pose a great threat to public health and the economy worldwide. Reports suggest their prevalence is increasing and can no longer be ignored. A detailed account on the current nature of the problem and identification of knowledge limitations in terms of geographical location, medicine classes, and type of medicine analysis performed is not available. Our objective was to systematically review articles that have reported investigations of counterfeit and substandard medicines.

Design: Systematic review.


Data Selection: Prospective field quality surveys on counterfeit and substandard medicines were selected from all available records within the selected databases up to December 31, 2013. All prospective studies performing chemical analysis on medicine samples were identified using the key search terms “counterfeit” or “substandard” and “medicine” or “drug” or “pharmaceutical.” The title, abstract, and/or full articles were reviewed for relevance according to a predetermined set of inclusion and exclusion criteria. Medicines procured from the Internet are beyond the scope of this review.

Results: Sixty-six research articles were found that fulfilled our inclusion criteria. The majority of medicine quality surveys were conducted in specific areas of Africa and Asia. Within these two continents, medicine quality reports covering the Northern part of Africa and the Western part of Asia in the Middle East are extremely scarce. Other continents such as North or South America and Europe were covered in limited articles, whereas the Australian continent had no reports. Moreover, most studies examined medicines that treat infectious diseases; very few articles addressed popular medicines for chronic diseases or clinically significant narrow therapeutic index medicines or cancer treatments, despite media reports of quality problems in these medicines. Furthermore, only six (9%) research articles attempted all levels of medicine quality analysis available through laboratory analysis, authentication of source, and package inspection to comprehensively identify the nature of the problem and so conclude whether the medicines were counterfeit or substandard.

Conclusion: Substandard and counterfeit medicines should be considered and identified through means of chemical analysis, physical analysis, authentication of source, and package inspection in any field medicine quality survey. More research is encouraged to examine the medicine quality in neglected parts of the globe and on neglected, yet popular and clinically significant, noncommunicable disease medicines.

Keywords: counterfeit, substandard, poor quality, SSFFC, medicine and drug

Introduction: Medicine safety, efficacy, and quality are the most important criteria in ensuring optimal treatment from medicines and are currently receiving increased attention in an era of
globalization and generic manufacturing.\textsuperscript{1,2} Medicines with questionable quality could either be counterfeit or substandard, according to the World Health Organization (WHO). A counterfeit medicine is defined by the WHO as “one which is deliberately and fraudulently mislabeled with respect to identity and/or source.” Counterfeiting could include both branded and generic products and may include products with the correct ingredients or with the wrong ingredients, without active ingredients, with insufficient active ingredient, or with fake packaging.\textsuperscript{3} Substandard medicines, also referred to as out-of-specification products, are defined by the WHO as “products that do not meet the required specification in terms of content and ingredients.”\textsuperscript{4,5} They are legally manufactured but do not conform to specifications as a result of inadequate manufacturing or poor storage conditions.\textsuperscript{6–9} Recently, the term substandard/spurious/falsely labeled/falsified/counterfeit medicines (SSFFC) was used by the WHO to simultaneously describe both counterfeit and substandard medicines.\textsuperscript{10} This joint definition highlights the importance of identifying both counterfeit and substandard medicines in any proposed medicine quality survey.

The distinction between counterfeit and substandard medicines is imperative when applying appropriate strategies to combat potential threats of either quality problem.\textsuperscript{11,12} However, some dismiss this notion and argue that both counterfeit and substandard medicines are similar because they both claim to be something that in reality they are not.\textsuperscript{13} Nevertheless, correctly identifying the type of medicine quality problem could aid governments and responsible bodies in determining the need to involve local or international law enforcement, particularly when scarce economic resources are present. Counterfeit medicines are strongly linked with organized crime and would most likely require criminal experts to aid health care professionals to combat this problem, as demonstrated by the establishment of the International Medical Products Anti-Counterfeiting Taskforce to support the WHO efforts to combat counterfeit medicines globally.\textsuperscript{14}

Medicine quality problems could be fatal in extreme clinical outcomes and have also been associated with severe economic consequences. More than 700,000 deaths from tuberculosis and malaria have been strongly linked with ineffective counterfeit and substandard medicines worldwide.\textsuperscript{15,16} Mortality has also been reported after heparin contamination in the United States and sexual enhancement drugs adulterated with large contents of hypoglycemic drugs in Singapore.\textsuperscript{17–20} Moreover, substandard and counterfeit medicines have been related to morbidity, drug resistance, therapeutic failure, and toxicity.\textsuperscript{8,13,15,16} Economically, substandard and counterfeit medicines have been suggested to cause macroeconomic burdens worldwide by wasting limited resources, causing loss of productivity, and limiting investment of major pharmaceutical companies into medicine research and development.\textsuperscript{7,8,21} Furthermore, consequences of substandard and counterfeit medicines could result in loss of confidence in health care professionals and/or services.\textsuperscript{8,13,15,16}

The WHO estimates that around 10% of all global pharmaceutical supply is counterfeit and substandard, reaching up to 50% of the supply in developing countries and as low as 1% in the developed world.\textsuperscript{6,15,22} Moreover, it has been suggested that the majority of reported SSFFC medicines were substandard, rather than counterfeit, yet they receive far less attention within the media and the scientific community.\textsuperscript{23,24} Determining the exact prevalence rates of either counterfeit or substandard medicines could be a complex task and requires high-quality country-based medicine surveys, which are limited within the available literature.

The aim of this systematic review is to broadly explore the evidence of substandard and counterfeit medicines in scientific reports to identify current knowledge limitations and provide an overview report of the current situation. Previously, some reviews have focused on specific medicine categories or problems.\textsuperscript{13,23,25,26} Only one review comprehensively searched for substandard and counterfeit medicine articles covering the period from 1966 to 2006 without specifying a therapeutic medicine category.\textsuperscript{27} Recently, the first systematic review on the subject of counterfeit and substandard medicines was published.\textsuperscript{28} However, Almuzaini et al have only reviewed some articles from a single therapeutic class that demonstrated high-quality reporting, which could be useful in the determination of SSFFC prevalence rates but may not be comprehensive enough to describe the broad scope and nature of SSFFC medicines available in other reports. Further, the previous systematic review did not discuss the types of analysis performed in the included studies, nor did it identify therapeutic classes or global regions in which the quality of medicines remains largely unknown. This review attempts to cover these issues broadly to encourage future researchers on medicine quality to focus their attention on neglected medicines and neglected parts of the globe. Furthermore, this review discusses types of analysis currently performed in medicine quality surveys to identify areas of concern and to promote the consideration of counterfeit as well as substandard medicines when conducting any medicine quality survey.
Methods
Searching the literature
Scopus, PubMed, and ISI Web of Knowledge databases have been searched for relevant research articles. The search covered the period from 1997, the year the first relevant citation was found, up to December 31, 2013. There was no language restriction applied on our search results.

The following key search terms were used in conjunction, using (AND) to identify related articles: substandard(s) or counterfeit(s); medicine(s) or drug(s) or pharmaceutical(s). The choice of key search terms was based on key search terms used in five previous literature reviews. The main distinction of our present review compared with most previously published reviews is its systematic nature and broader scope, as no medicine groups or settings were specifically chosen in the search terms and inclusion criteria used.

The definitions and criteria used to describe counterfeit and substandard medicines in this review are based on the widely accepted WHO definitions of each phenomenon, as cited earlier. On the basis of the WHO criteria, a counterfeit medicine could be determined by chemical analysis methods if medicine samples contained no, or the wrong, active ingredient. A counterfeit medicine could also be identified via medicine package analysis by visual comparison to a known genuine package. Other means of detecting counterfeit medicines include authenticating its source through official consignment documents or communication with the stated manufacturer and regulatory organizations. In addition, deliberately manufactured substandard medicines are considered counterfeit, although this would be difficult to demonstrate without legal and criminal investigation by authorities. In contrast, a substandard medicine should always contain the correct active pharmaceutical ingredient (API), be produced from a legitimate source, and be without packaging defaults. Substandard medicines are present when the amount of API is outside the acceptable pharmacopeial limits, the sample does not meet other standards set by the pharmacopoeias, or medicines are past their expiry dates. Collectively, we refer to both counterfeit and substandard medicines as SSFFC medicines, in accordance with the latest WHO joint definition.

Inclusion and exclusion criteria for articles in this review
Studies included in this review were original research articles that reported prospective medicine sample collection from their natural settings; these medicines were presumed to be readily available to patients. Further, all included articles must have reported conducting chemical tests for the identification and/or quantification of the API. Without performing chemical analysis, it would not be possible to determine whether a medicine sample was counterfeit or not, as no information on the API would be present. In addition, relevant studies would include medicine samples from a wide range of different therapeutic categories and dosage forms without any restrictions.

In contrast, the exclusion criteria of articles would include studies that did not report primary collection of medicine samples or medicines procured from the Internet or retrospectively collected through authority or innovator company seizures. Furthermore, studies that reported only physical or packaging testing without chemical analysis were excluded. Duplicate results and nonrelevant articles were also identified and excluded from this review.

Data presentation of articles in this review
This systematic review has been performed in accordance with Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines for systematic reviews. All percentages of SSFFC medicines available in this review are reported as cited from their primary source. Therefore, caution is advised, as methodological differences exist between articles. The data presented here do not allow for any estimation of the SSFFC prevalence rate worldwide.

Results
Data extraction
The use of the selected search terms resulted in a total of 3,861 hits from all databases. An initial screening of titles/abstracts followed this, excluding nonrelevant and duplicate results to reduce the number of results to 1,288 research articles. Subsequently, a full review of articles was performed that further excluded articles without primary data collection, such as reviews and opinions, articles containing retrospective sample collection of medicines (either donated or seized by authorities), medicines acquired through the Internet, nonrelated articles, studies without medicine sample collection, and studies that did not perform chemical analysis of samples. This strategy reduced the final number of the included articles to 66. A flowchart illustrating the method used for article selection in this review and different exclusion categories is shown in Figure 1.
The majority of reported studies prospectively examining SSFFC medicines were conducted in the African continent (31/66; 47%). Nigeria and Ghana alone were selected for more than 50% (17/31) of the studies in Africa. In Asia, 23/66 (35%) of the SSFFC medicine quality surveys were conducted, mostly in the South Eastern part of Asia (Tables 1–4). Eight research articles were performed in the southern parts of the continent in Pakistan, Bangladesh, and India. Overall, only two studies (3%) were published that addressed SSFFC medicines in the western part of Asia, also known to be part of the Middle East. Elsewhere, 6/66 (9%) of studies were conducted in more than one continent simultaneously. Moreover, three studies were performed in North/South America (4%) and two in Eastern Europe (3%). Only one study was located in the borderline area between Asia and Australia in Papua New Guinea.

**Medicine therapeutic classes**

Substandard and counterfeit medicines were found from various therapeutic categories. However, most SSFFC studies (57/66; 86%) were focused on medicines that treat infectious diseases. Antimalarial, antibiotic, and antituberculosis medicines were examined in 30/66 (46%), 10/66 (15%), and 5/66 (8%) of the located studies, respectively (Tables 1, 2, and 4). The combination of more than one class of medicines to treat infectious diseases was found in 12/66 (18%) of the articles. Other infectious diseases such as leishmaniasis medicines were investigated on one (2%) other occasion. In contrast, medicines for treatment of non-communicable diseases were present in only 9/66 (14%) of the cited literature. The analgesic paracetamol was investigated on two separate occasions. Similarly, antihypertensive medications were surveyed in only two studies. Nonsteroidal anti-inflammatory agent aspirin was analyzed in one further study. The antihistamine medicine chlorpheniramine was only present in one survey. Narrow-therapeutic index medicines also were the focus of only one published study. Other types of medicines such as ergometrine, oxytocin, and erythropoietin appeared in only one study each. A single study attempted to collect samples from various therapeutic categories simultaneously.

**Evidence and nature of SSFFC medicines**

Overall, substandard medicines were found in the majority of prospective SSFFC medicine studies (60/66; 91%) (Tables 1 and 4). Counterfeit medicines were less evident in
### Table 1 Research articles reporting both counterfeit and substandard medicines

<table>
<thead>
<tr>
<th>Reference</th>
<th>Country</th>
<th>Medicine</th>
<th>Sample size</th>
<th>Authenticate source</th>
<th>Visual analysis</th>
<th>Chemical analysis</th>
<th>Physical analysis</th>
<th>Results</th>
<th>Type of substandard/spurious/false labeled/falsified/counterfeit medicine problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bate et al.</td>
<td>17 countries from all continents</td>
<td>Antituberculosis isoniazid and rifampicin</td>
<td>713</td>
<td>NR</td>
<td>Package</td>
<td>TLC</td>
<td>Disintegration</td>
<td>65/713 (9.1%) substandard; 18/713 (2.5%) counterfeit</td>
<td>Low API%; no API or suspicious packaging</td>
</tr>
<tr>
<td>Stanton et al.</td>
<td>Ghana</td>
<td>Ergometrine, oxytocin</td>
<td>101</td>
<td>NR</td>
<td>NR</td>
<td>Performed by local food and drug administrations</td>
<td>NR</td>
<td>92/101 (91%) substandard; 1/101 (1%) counterfeit</td>
<td>Low or high API%, 2 expired; no API</td>
</tr>
<tr>
<td>Baratta et al.</td>
<td>15 different countries</td>
<td>Various therapeutic classes and formulations</td>
<td>196</td>
<td>NR</td>
<td>NR</td>
<td>UV and some with HPLC</td>
<td>Uniformity of content, mass, disintegration, friability and hardness tests</td>
<td>101/196 (52%) substandard; 4/196 (2%) counterfeit</td>
<td>Various failures mostly physical; no API</td>
</tr>
<tr>
<td>Nair et al.</td>
<td>Papua New Guinea</td>
<td>Antimalarial amodiaquine and antibiotic amoxicillin</td>
<td>14</td>
<td>Internet search and e-mail contact manufacturer</td>
<td>Package and label inspection</td>
<td>TLC and HPLC</td>
<td>Weight variation, content uniformity, dissolution</td>
<td>11/14 (79%) substandard; 3/14 (11%) counterfeit</td>
<td>Poor content uniformity, fails assay and inappropriate packaging; no API, no manufacturer address, and distributor does not exist</td>
</tr>
<tr>
<td>Ali et al.</td>
<td>Nigeria</td>
<td>Antimalarial ACT</td>
<td>6</td>
<td>NR</td>
<td>Package inspection</td>
<td>UV</td>
<td>NR</td>
<td>3/6 (50%) substandard; 2/6 (33%) counterfeit</td>
<td>Low API%; missing manufacturer details on package, and no expiry date</td>
</tr>
<tr>
<td>Khan et al.</td>
<td>Cambodia</td>
<td>Albendazole, mebendazole, and metronidazole</td>
<td>203</td>
<td>Contact with manufacturer and authorities</td>
<td>Package inspection</td>
<td>HPLC</td>
<td>Disintegration and weight measurement</td>
<td>2% substandard; 4% counterfeit</td>
<td>Failed disintegration; failed authenticity with manufacturer or authorities</td>
</tr>
<tr>
<td>Ochekpe et al.</td>
<td>Nigeria</td>
<td>Antimalarial artemisinin combination therapies</td>
<td>70</td>
<td>NR</td>
<td>Package inspection Minilab® (Global Pharma Health Fund eV, Giessen, Germany)</td>
<td>TLC</td>
<td>Disintegration</td>
<td>27/70 (38%) substandard; 4/70 (6%) counterfeit</td>
<td>Low API%; no API and fake packaging</td>
</tr>
<tr>
<td>Bate et al.</td>
<td>India</td>
<td>Antimalarial, antibiotic, and antitubercular</td>
<td>541</td>
<td>NR</td>
<td>NR</td>
<td>TLC</td>
<td>Disintegration</td>
<td>46/541 (8.5%) substandard; 11/541 (2%) counterfeit</td>
<td>Low API% and disintegration failure; no API</td>
</tr>
<tr>
<td>Onwujekwe et al.</td>
<td>Nigeria</td>
<td>Antimalarial</td>
<td>225</td>
<td>NR</td>
<td>NR</td>
<td>HPLC</td>
<td>Dissolution</td>
<td>60/225 (37%) substandard or counterfeit</td>
<td>Less API% or wrong API</td>
</tr>
<tr>
<td>Risha et al.</td>
<td>Tanzania</td>
<td>Antimalarial, antibiotic, and antiretroviral</td>
<td>1,257</td>
<td>NR</td>
<td>Package inspection Minilab® and TLC</td>
<td>Color reaction test and dissolution</td>
<td>46/1,257 (3.6%) substandard; 5/1,257 (0.4%) counterfeit</td>
<td>Dissolution failure mostly; NR</td>
<td></td>
</tr>
<tr>
<td>Tipke et al.</td>
<td>Burkina Faso</td>
<td>Antimalarial</td>
<td>77</td>
<td>Internet search was for manufacturers</td>
<td>Package inspection Minilab® and TLC</td>
<td>Color reaction and TLC</td>
<td>32/77 (42%) substandard; 47/77 (1.2%) counterfeit</td>
<td>Failed visual inspection, low API%, and failure of dissolution test; no API</td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Reference</th>
<th>Country</th>
<th>Medicine</th>
<th>Sample size</th>
<th>Authenticate source</th>
<th>Visual analysis</th>
<th>Chemical analysis</th>
<th>Physical analysis</th>
<th>Results</th>
<th>Type of substandard/spurious/false labeled/falsified/counterfeit medicine problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bate et al</td>
<td>Six African countries</td>
<td>Antimalarial</td>
<td>210</td>
<td>NR</td>
<td>Package inspection</td>
<td>TLC</td>
<td>Dissolution</td>
<td>35% (73/210) substandard; 7/210 (3%) counterfeit</td>
<td>Low API% and dissolution failure; missing manufacturing and/or expiry date on the package</td>
</tr>
<tr>
<td>Pouillot et al</td>
<td>Cameroon and Niger</td>
<td>Antimalarial, antibiotic, and antihelminic</td>
<td>153</td>
<td>NR</td>
<td>Basic package inspection</td>
<td>HPLC and UV</td>
<td>Average weight and uniformity of mass, disintegration, dissolution</td>
<td>66/153 (43%) substandard; 5/153 (3%) counterfeit</td>
<td>Nonconforming to API% and physical tests; no API</td>
</tr>
<tr>
<td>Ofori-Kwakye et al</td>
<td>Ghana</td>
<td>Antimalarial artesunate</td>
<td>17</td>
<td>NR</td>
<td>Basic package inspection</td>
<td>Colorimetric and spectrometry</td>
<td>Uniformity of weight, breaking strength, friability, and rate of disintegration</td>
<td>11/17 (65%) substandard; 1/17 (6%) counterfeit</td>
<td>Failed content uniformity test; manufacturer address missing</td>
</tr>
<tr>
<td>Atemnkeng et al</td>
<td>Congo</td>
<td>Antimalarial</td>
<td>28</td>
<td>NR</td>
<td>Package inspection</td>
<td>UV, TLC, and HPLC-UV</td>
<td>NR</td>
<td>4/28 (14%) substandard; 13/28 (46%) counterfeit</td>
<td>Low and high API%; no API, no manufacturer name, and no trade name</td>
</tr>
<tr>
<td>Gaudiano et al</td>
<td>Congo, Burundi, and Angola</td>
<td>Antimalarial</td>
<td>30</td>
<td>NR</td>
<td>Package and label</td>
<td>HPLC</td>
<td>Uniformity of mass, disintegration, and dissolution</td>
<td>17/30 (57%) substandard; 1/30 (3%) counterfeit; 1/30 (3%) diverted</td>
<td>Low API% and physical test failures; no API; humanitarian medicine</td>
</tr>
<tr>
<td>Atemnkeng et al</td>
<td>Kenya and Congo</td>
<td>Antimalarial artemisinin-derivative drugs</td>
<td>24</td>
<td>Checked source of some companies</td>
<td>Check for illegal prints only</td>
<td>HPLC-UV</td>
<td>NR</td>
<td>9/24 (38%) substandard; 3/24 (12.5%) counterfeit</td>
<td>Low and high API%; nonexistent manufacturer</td>
</tr>
<tr>
<td>Syhakhang et al</td>
<td>Laos (two studies in 1997 and 1999)</td>
<td>Antibiotic, antimalarial, and aspirin</td>
<td>666</td>
<td>NR</td>
<td>Check for illegal prints only</td>
<td>HPLC, titration, and UV</td>
<td>Weight variation and disintegration</td>
<td>46% and 22% substandard in 1997 and 1999; 1% counterfeit</td>
<td>Low or high API% and failed weight variation; no API</td>
</tr>
<tr>
<td>Basco</td>
<td>Cameroon</td>
<td>Antimalarial</td>
<td>284</td>
<td>NR</td>
<td>NR</td>
<td>Color test and TLC</td>
<td>NR</td>
<td>53/284 (18%) substandard; 59/284 (20%) counterfeit</td>
<td>Low API%; no API</td>
</tr>
<tr>
<td>Dondorp et al</td>
<td>Thailand, Vietnam, Cambodia, Lao People's Democratic Republic, and Myanmar</td>
<td>Antimalarial artesunate derivatives and mefloquine</td>
<td>303</td>
<td>NR</td>
<td>Package analysis of holograms</td>
<td>Color test and HPLC</td>
<td>NR</td>
<td>49/303 (16%) counterfeet; 96/303 (32%) substandard</td>
<td>No or trace API, and all were artesunate; standard API% and all were mefloquine</td>
</tr>
<tr>
<td>Prazuck et al</td>
<td>Myanmar</td>
<td>Antibiotics</td>
<td>21</td>
<td>NR</td>
<td>UV, TLC, and titrimetry</td>
<td>NR</td>
<td>10/21 (48%) substandard; 3/21 (14%) counterfeit</td>
<td>Low, high API% and expired medicines; wrong API and no expiry date on package</td>
<td></td>
</tr>
</tbody>
</table>

Table 1 (Continued)
Counterfeit and substandard medicines in field quality surveys

Taylor et al. 51 Nigeria Antimalarial, antibiotic, antituberculosis, antifungal

581 NR Basic package information on origin

279/581 (48%) substandard; 43/581 (7%) counterfeit

Stenson et al. 52 Laos Antibiotics

366 NR Basic visual analysis

43/366 (11.5%) substandard; 12/366 (3.3%) counterfeit

Shakoor et al. 53 Nigeria and Thailand Antimalarial and antibiotics

96 NR Package inspection for obvious errors

36% from Nigeria and 40% from Thailand substandard; 6/96 (6%) counterfeit

Abbreviations: API, active pharmaceutical ingredient; NR, not reported; TLC, thin layer chromatography; HPLC, high-performance liquid chromatography; UV, ultraviolet; ACT, artemisinin combination therapies.

29/66 (44%) of available studies (Tables 1 and 2). Counterfeit and substandard medicines were simultaneously found in 24/66 (36%) articles (Table 1). Few studies 5/66 (8%) reported only evidence of counterfeiting in the medicine samples collected (Table 2). Evidence of medicines being only substandard, rather than counterfeit, was found in 36/66 (55%) of the articles (Table 4). One study did not find evidence of counterfeit or substandard medicines in their sample (Table 3).

Several types of SSFFC problems have been reported in the selected literature. It was noted that more than one medicine quality problem typically exists within each prospective medicine quality survey (Tables 1, 2, and 4). The most reported medicine quality problem was failure to comply with the specified API limits in 46/66 (70%) of cases (Tables 1 and 4). Failure of dissolution or disintegration tests has been reported in 24/66 (36%) of the articles (Tables 1 and 4). The presence of either no API12,31–33,36,42,44,45,47–49,51–56 or the wrong API12,38,50,55 was reported in 24/66 (30%) and 4/66 (6%) cases, respectively. Other problems were also reported, including fake package,36,57 fake hologram,12,56,57 manufacturer does not exist,12,33,46 manufacturer confirmed a nonauthentic batch,35,56 expired medicines,31,50,68 no origin country stated,51 no manufacturer address,33,34,43 no manufacturer stated,44 no expiry date,34,41,50 unusual interval between manufacturing and expiry date,55 wrong name on package or leaflet,12 wrong spelling of “tablet,”55,56 use of a different font,56 different medicinal taste,57 heavier weight,57 unauthorized manufacturer,66 absence of trade name,44 signs of deterioration,53 and diverted medicines45,80 intended for distribution in one location and found to be on sale in another market.

Type of analysis identified in the included studies

Four distinctive types of analysis can be used to distinguish between a genuine and SSFFC medicines; namely, authentication of the supplier, visual package inspection, and chemical and physical analysis (Tables 1–4). Authentication of the medicine source via contact with manufacturer, health regulatory agencies, or Internet search has been only attempted in 10/66 (15%) of the selected studies.12,33,35,40,46,56,62,80,86,88 Package inspection was more popular than authentication, being reported in 39/66 (59%) of studies, with the majority reporting obvious spelling errors and basic label information (medicine name, dosage, manufacturer, expiry date, and lot number), as shown in Tables 1, 2, and 4. As for the chemical analysis, high-performance liquid chromatography and thin-layer chromatography (TLC) were most widely used in 40/66
## Table 2 Research articles reporting counterfeit medicines only

<table>
<thead>
<tr>
<th>Reference</th>
<th>Country</th>
<th>Medicine(s)</th>
<th>Sample size</th>
<th>Authenticate source</th>
<th>Visual analysis</th>
<th>Chemical analysis</th>
<th>Physical analysis</th>
<th>Results</th>
<th>Type of substandard/spurious/falsely labeled/falsified/counterfeit medicine problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dorlo et al 14</td>
<td>Bangladesh</td>
<td>Miltefosine</td>
<td>2</td>
<td>NR</td>
<td>Package inspection</td>
<td>Liquid chromatography-mass spectrometry, Fourier transform infrared spectroscopy, near-infrared spectroscopy, colorimetric test</td>
<td>NR</td>
<td>Both (100%) failed all tests and are counterfeit</td>
<td>No API</td>
</tr>
<tr>
<td>Newton et al 12</td>
<td>Multiple countries in Africa</td>
<td>Antimalarial</td>
<td>59</td>
<td>Contact manufacturer</td>
<td>Package inspection</td>
<td>HPLC, mass spectroscopy, pollen analysis, X-ray diffraction</td>
<td>NR</td>
<td>Only case reports of counterfeits and do not allow for percentage estimation</td>
<td>Wrong API, nonexistent manufacturer, no API, hologram different from genuine package, and wrong name on packaging or leaflet</td>
</tr>
<tr>
<td>Sengaloundeth et al 15</td>
<td>Laos</td>
<td>Antimalarial artesunate</td>
<td>30</td>
<td>NR</td>
<td>Package analysis</td>
<td>Colorimetric tests, HPLC, mass spectroscopy, pollen analysis, X-ray diffraction</td>
<td>NR</td>
<td>88% failure and counterfeit</td>
<td>No or wrong API, wrong spelling of “tablet” on package and unusual interval between manufacturing date and expiry date of 9 years</td>
</tr>
<tr>
<td>Newton et al 16</td>
<td>Vietnam, Cambodia, Myanmar, Laos, and Thailand</td>
<td>Antimalarial artesunate</td>
<td>391</td>
<td>Contact with one company to authenticate batch numbers</td>
<td>Package analysis including holograms</td>
<td>Colorimetric, HPLC, mass spectrometry</td>
<td>NR</td>
<td>195/391 (50%) counterfeit</td>
<td>Fake hologram, wrong spelling on packaging, use of different font, failure of authentication when manufacturer was contacted, and no API</td>
</tr>
<tr>
<td>Newton et al 17</td>
<td>Myanmar, Cambodia, Vietnam, Laos, and western Thailand</td>
<td>Antimalarial artesunate</td>
<td>104</td>
<td>NR</td>
<td>Package inspection and holograms, printing, and bar codes</td>
<td>Color reaction test</td>
<td>Tablet weight, size, and color</td>
<td>Overall, 38% are counterfeit found in all countries</td>
<td>No API, different taste of tablets, heavier weight of tablets, and different packaging and holograms compared with genuine</td>
</tr>
</tbody>
</table>

**Abbreviations:** API, active pharmaceutical ingredient; NR, not reported; HPLC, high-performance liquid chromatography.
(61%) and 19/66 (29%) of studies, respectively (Tables 1, 2, and 4). Other chemical analysis methods were reported such as color reaction tests,\textsuperscript{32,42,43,45,46,69} spectroscopic techniques,\textsuperscript{12,43,54–56,67,73} and titration,\textsuperscript{47} but remain less frequently used. Moreover, physical analysis tests were performed in 39/66 (59%) of the studies (Tables 1, 2 and 4). The most common physical tests reported were disintegration and/or dissolution tests in 36/39 (92%) cases (Tables 1, 2, and 4). Other less frequently used physical analysis tests include content uniformity,\textsuperscript{33,42,43,45,68} weight measurement,\textsuperscript{33,35,42,47,52,57,60,63,65,67,73} hardness,\textsuperscript{32,65,73} and friability\textsuperscript{32,43,60,65,73} tests. Interestingly, only six studies (9%) reported all four types of analysis in an attempt to clearly identify and classify the type of SSFFC problem, where present, in any medicine sample.\textsuperscript{33,35,40,62,86,88}

### Discussion

#### Neglected parts of the world in SSFFC surveys

According to our findings, the vast majority of prospective medicine quality studies were conducted in small parts of Africa and Asia. These efforts can be attributed to an attempt to counteract nonexistent or lower levels of regulation in these pharmaceutical markets.\textsuperscript{94} However, some parts of these two continents still have limited scientific research addressing the problem of SSFFC medicines, mainly in the Middle East and North Africa. In Yemen, 32% of selected antimalarial medicines failed analysis tests, and the majority of these were substandard, having lower than accepted API% limits and unacceptable dissolution rates.\textsuperscript{87} Another study explored the API content of the antibiotic amoxicillin purchased from Egypt, Lebanon, Jordan, and Saudi Arabia and found that more than 50% of samples had lower API% than accepted by pharmacopeial limits, and therefore were considered substandard.\textsuperscript{82} A multicountry medicine quality survey found that 12% of samples collected from Egypt failed at least one medicine quality test and can be considered substandard.\textsuperscript{71} None of these studies reported an attempt to verify the source or analyze packages of the selected medicine samples to explore the possibility of counterfeiting activity. This may cause some concern, particularly with recent seizures of SSFFC medicines in this area. In addition, the currently unsettled political situation may be a catalyst for the increased prevalence of SSFFC medicines, as it allows them to escape immediate governmental attention.\textsuperscript{95} Reports of recent seizures of SSFFC medicines in this area can be mostly found in the media, which remains the main source of information regarding SSFFC medicines in this region with limited published scientific reports.\textsuperscript{95} Moreover, a WHO report on questionnaire responses from a number of health organizations in the Eastern Mediterranean Regional Office regarding counterfeit medicines has confirmed counterfeit seizures in this region by some respondent countries.\textsuperscript{96} In addition, this area could be of specific importance in terms of geographical location, as it separates two well-established regions of SSFFC medicine prevalence, according to our data, and is en route between potential counterfeit manufacturers in Asia\textsuperscript{88} and their global targeted markets. It is therefore suggested that several pilot studies be conducted to survey the quality of medicines in the Middle East and North Africa to assess the current medicine quality situation before any countermeasures or large-scale medicine quality surveys can be recommended. Elsewhere, such pilot studies have been shown to be instrumental in the assessment of the medicine quality situation in different countries and to have justified the need for further medicine quality surveys, where appropriate.\textsuperscript{30,37,57,78}

Evidence from South America suggests that SSFFC medicines are available, but with only limited scientific research. A study found 11% of antimalarials to be substandard in seven South American countries using basic TLC chemical analysis.\textsuperscript{84} The TLC analysis technique is limited by its inability to detect higher than 80% of API concentration in medicine samples\textsuperscript{41} which has been evident to exist in previous studies.\textsuperscript{31,41,44,46,50,52,53,61,63,66,67,73,76–78,87,91,93} It is therefore possible that the prevalence of SSFFC medicines in South America could be higher than the reported figures if more sophisticated chemical techniques for the quantification of API% content were used, such as high-performance liquid chromatography. Another study reported problems with low API% on a range of medicines procured from Mexico; of particular importance are some narrow therapeutic index

### Table 3 Study with no report of substandard or counterfeit medicines

<table>
<thead>
<tr>
<th>Reference</th>
<th>Country</th>
<th>Medicine</th>
<th>Sample size</th>
<th>Authenticate source</th>
<th>Visual analysis</th>
<th>Chemical analysis</th>
<th>Physical analysis</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Said et al\textsuperscript{88}</td>
<td>Malaysia</td>
<td>Paracetamol</td>
<td>16</td>
<td>NR</td>
<td>NR</td>
<td>Near-infrared spectroscopy</td>
<td>NR</td>
<td>All samples passed but with variable quality</td>
</tr>
</tbody>
</table>

Abbreviation: NR, not reported.
### Table 4: Research articles reporting only substandard medicines

<table>
<thead>
<tr>
<th>Reference</th>
<th>Country</th>
<th>Medicine</th>
<th>Sample size</th>
<th>Authenticate source</th>
<th>Visual analysis</th>
<th>Chemical analysis</th>
<th>Physical analysis</th>
<th>Results</th>
<th>Type of substandard/spurious/falsely labeled/falsified/counterfeit medicine problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haruna et al19</td>
<td>Nigeria</td>
<td>Antihypertensive methyldopa</td>
<td>4</td>
<td>NR</td>
<td>NR</td>
<td>Nonaqueous titration</td>
<td>NR</td>
<td>1/4 (25%)</td>
<td>Low API%</td>
</tr>
<tr>
<td>Audu et al20</td>
<td>Congo</td>
<td>Antihistamine chlorpheniramine</td>
<td>10</td>
<td>NR</td>
<td>NR</td>
<td>HPLC and UV</td>
<td>Tablet shape, size, thickness, and weight; disintegration and friability tests</td>
<td>3/10 (30%)</td>
<td>Low API%</td>
</tr>
<tr>
<td>Ramachandran et al21</td>
<td>India</td>
<td>Anti-TB</td>
<td>1,948 tablets</td>
<td>NR</td>
<td>NR</td>
<td>Spectrometry</td>
<td>NR</td>
<td>168/1,948 (9%)</td>
<td>Low and high API%</td>
</tr>
<tr>
<td>Khan et al22</td>
<td>Cambodia</td>
<td>Antibiotic amoxicillin-clavulanic acid</td>
<td>59</td>
<td>Contact with manufacturer and local authorities</td>
<td>Basic visual analysis of primary and secondary packaging</td>
<td>HPLC</td>
<td>Stability and dissolution</td>
<td>12/59 (20%)</td>
<td>Low API%, failure of content uniformity and dissolution tests</td>
</tr>
<tr>
<td>Affum et al23</td>
<td>Ghana</td>
<td>Antimalarial artesunate and amodiaquine</td>
<td>32 blisters</td>
<td>NR</td>
<td>Basic visual analysis compared to genuine</td>
<td>Titrimetric, HPLC, and spectrometry</td>
<td>Tablet weight</td>
<td>14/32 (43.75%)</td>
<td>Low and high API% mostly artesunate</td>
</tr>
<tr>
<td>Briesen et al24</td>
<td>Kenya and Congo</td>
<td>Antimicrobial eye drops</td>
<td>33</td>
<td>NR</td>
<td>NR</td>
<td>HPLC</td>
<td>NR</td>
<td>19/33 (58%)</td>
<td>Low and high API%</td>
</tr>
<tr>
<td>Nogueira et al25</td>
<td>Brazil</td>
<td>Antimalarial medicines</td>
<td>9</td>
<td>NR</td>
<td>Simple package analysis</td>
<td>HPLC-UV</td>
<td>Dissolution, disintegration, hardness, uniformity of weight, and friability tests</td>
<td>4/9 (44%)</td>
<td>Failing only visual inspection and uniformity of weight</td>
</tr>
<tr>
<td>El-Duah and Ofori-Kwakye26</td>
<td>Ghana</td>
<td>Antimalarial artemisinin-medicines</td>
<td>14</td>
<td>NR</td>
<td>For illegal print errors</td>
<td>Colorimetry and TLC</td>
<td>Uniformity of mass, crushing strength, and disintegration</td>
<td>13/14 (93%)</td>
<td>Low or high API% and failing physical tests</td>
</tr>
<tr>
<td>Karlage et al27</td>
<td>Mexico</td>
<td>Antibiotics, warfarin, levothyroxine and sildenafil</td>
<td>17</td>
<td>NR</td>
<td>HPLC</td>
<td>Weight measurement</td>
<td>5/17 (30%)</td>
<td>Low API%</td>
<td></td>
</tr>
<tr>
<td>Pribluda et al28</td>
<td>Seven countries in South America</td>
<td>Antimalarial</td>
<td>1,663</td>
<td>NR</td>
<td>Package and label</td>
<td>TLC</td>
<td>Disintegration</td>
<td>193/1,663 (11%)</td>
<td>Expired medicines mostly, low API%, and failure of disintegration tests</td>
</tr>
<tr>
<td>Klein et al29</td>
<td>Ghana</td>
<td>Antimalarial</td>
<td>33</td>
<td>NR</td>
<td>Package inspection</td>
<td>NMR</td>
<td>NR</td>
<td>1/33 (3%)</td>
<td>Low API%</td>
</tr>
<tr>
<td>Ehianeta et al30</td>
<td>Nigeria</td>
<td>Antimalarial artesunate and amodiaquine combination</td>
<td>13</td>
<td>NR</td>
<td>Package inspection of expiry date and registration</td>
<td>HPLC</td>
<td>NR</td>
<td>11/13 (85%)</td>
<td>Low and high API%</td>
</tr>
<tr>
<td>Authors</td>
<td>Countries</td>
<td>Drugs Description</td>
<td>Number of Samples</td>
<td>Methods of Analysis</td>
<td>Test Results</td>
<td>Substandard (%)</td>
<td>Reason for Failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------------</td>
<td>----------------------------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Bate et al²¹</td>
<td>17 countries from all continents</td>
<td>Antimalarial, antibiotics, and antituberculosis</td>
<td>899</td>
<td>Package inspection</td>
<td>Disintegration</td>
<td>15%</td>
<td>Failure of visual inspection, low API%, and dissolution failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seear et al²²</td>
<td>India</td>
<td>Ciprofloxacin, artemesunate, and rifampicin</td>
<td>300</td>
<td>NR</td>
<td>HPLC-MS</td>
<td>43%</td>
<td>Low and high API%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Akpabio et al²³</td>
<td>Nigeria</td>
<td>Antibiotic ciprofloxacin</td>
<td>4</td>
<td>NR</td>
<td>Titration</td>
<td>1/4 (25%)</td>
<td>Low API%, failure of friability, and dissolution tests</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hadi et al²⁴</td>
<td>Indonesia</td>
<td>Five different antibiotics</td>
<td>104</td>
<td>Package inspection</td>
<td>HPLC</td>
<td>18%</td>
<td>Low API%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bate and Hess²⁵</td>
<td>Ghana and Nigeria</td>
<td>Antimalarial</td>
<td>339</td>
<td>Package inspection</td>
<td>Disintegration</td>
<td>23%</td>
<td>Failure of visual inspection, low API%, and dissolution failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leslie et al²⁶</td>
<td>Pakistan</td>
<td>Antimalarial</td>
<td>9</td>
<td>NR</td>
<td>HPLC</td>
<td>100%</td>
<td>High API% and dissolution failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Twagirumukiza et al²⁷</td>
<td>Rwanda</td>
<td>Antihypertensive drugs</td>
<td>10</td>
<td>NR</td>
<td>HPLC</td>
<td>2/10 (20%)</td>
<td>Low and high API%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ali²⁸</td>
<td>Pakistan</td>
<td>Antibiotic ceftriaxone injection</td>
<td>96</td>
<td>NR</td>
<td>HPLC</td>
<td>15/96 (16%)</td>
<td>Low and high API%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bate et al²⁹</td>
<td>Ghana, India, Kenya, Nigeria, Tanzania, and Uganda</td>
<td>Antimalarial, antibiotic, and antimycobacterial</td>
<td>78</td>
<td>NR</td>
<td>Disintegration</td>
<td>40/78 (51%)</td>
<td>Low API% and disintegration failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fotiou et al³⁰</td>
<td>Thailand</td>
<td>Epoetin alfa-pre-filled syringes</td>
<td>139</td>
<td>Checked batch numbers with manufacturer</td>
<td>Primary and secondary package and security features</td>
<td>HPLC, electrophoresis and Western blotting</td>
<td>32/139 (23%)</td>
<td>Exceeded specific content requirement for the product, and batch number matches products sold outside the country according to the manufacturer</td>
<td></td>
</tr>
<tr>
<td>Kaur et al³¹</td>
<td>Tanzania</td>
<td>Antimalarial</td>
<td>304</td>
<td>NR</td>
<td>HPLC</td>
<td>12.5%</td>
<td>Low API%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kyriacos et al³²</td>
<td>Lebanon, Syria, Jordan, Egypt, and Saudi Arabia</td>
<td>Amoxicillin antibiotic in different formulations</td>
<td>111</td>
<td>NR</td>
<td>HPLC</td>
<td>56%</td>
<td>Low API%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meos et al³³</td>
<td>Estonia and Russia</td>
<td>Antibiotic doxycycline</td>
<td>8</td>
<td>NR</td>
<td>HPLC</td>
<td>2/8 (25%)</td>
<td>Low API% and dissolution failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bronnikova et al³⁴</td>
<td>Estonia and Russia</td>
<td>Antibiotic amoxicillin</td>
<td>6</td>
<td>NR</td>
<td>HPLC and UV</td>
<td>1/6 (16%)</td>
<td>Dissolution failure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vijaykadga et al³⁵</td>
<td>Thailand</td>
<td>Antimalarial</td>
<td>369</td>
<td>Package and label</td>
<td>Disintegration</td>
<td>23/369 (6%)</td>
<td>Low API% and disintegration test failure</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
Table 4 (Continued)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Country</th>
<th>Medicine</th>
<th>Sample size</th>
<th>Authenticate source</th>
<th>Visual analysis</th>
<th>Chemical analysis</th>
<th>Physical analysis</th>
<th>Results</th>
<th>Type of substandard/spurious/falsey labeled/falsified/counterfeit medicine problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lon et al</td>
<td>Cambodia</td>
<td>Antimalarial</td>
<td>451</td>
<td>NR</td>
<td>Visual inspection</td>
<td>TLC</td>
<td>Disintegration</td>
<td>122/451 (27%)</td>
<td>Low API% and disintegration test failure; one illegal manufacturer identified</td>
</tr>
<tr>
<td>Amin et al</td>
<td>Kenya</td>
<td>Antimalarial</td>
<td>116</td>
<td>NR</td>
<td>Package and storage area inspection</td>
<td>UV and HPLC</td>
<td>Dissolution</td>
<td>47/116 (40%)</td>
<td>Low API% and dissolution failure</td>
</tr>
<tr>
<td>Abdoo-Rabbo et al</td>
<td>Yemen</td>
<td>Antimalarial tablet and syrup</td>
<td>50</td>
<td>NR</td>
<td>UV and HPLC</td>
<td>Dissolution</td>
<td>16/50 (32%)</td>
<td>Low API%, high API% and dissolution failure</td>
<td></td>
</tr>
<tr>
<td>Rookkapan et al</td>
<td>Thailand</td>
<td>Antituberculosis</td>
<td>52</td>
<td>One quality report was requested from a manufacturer</td>
<td>Tablet inspection</td>
<td>UV and HPLC</td>
<td>Dissolution</td>
<td>37% substandard</td>
<td>Failure of visual inspection, low API%, and dissolution failure</td>
</tr>
<tr>
<td>Kayumba et al</td>
<td>Rwanda and Tanzania</td>
<td>Antimicrobial and antimalarial drugs</td>
<td>33</td>
<td>NR</td>
<td>HPLC</td>
<td>Dissolution</td>
<td>4/33 (12%)</td>
<td>4/33 (12%)</td>
<td>Dissolution failure</td>
</tr>
<tr>
<td>Minzi et al</td>
<td>Tanzania</td>
<td>Antimalarial</td>
<td>33</td>
<td>NR</td>
<td>Basic package information</td>
<td>TLC, HPLC</td>
<td>Dissolution</td>
<td>12/33 (36%)</td>
<td>Low API% and dissolution failure</td>
</tr>
<tr>
<td>Obodzie et al</td>
<td>Nigeria</td>
<td>Antibiotic in different formulation</td>
<td>22</td>
<td>NR</td>
<td>HPLC</td>
<td>NR</td>
<td>9/22 (41%)</td>
<td>Low and high API%</td>
<td></td>
</tr>
<tr>
<td>Laserson et al</td>
<td>Seven different countries</td>
<td>Anti-TB</td>
<td>71</td>
<td>NR</td>
<td>Basic package information</td>
<td>TLC and LC-MS</td>
<td>NR</td>
<td>10% substandard</td>
<td>Low API%</td>
</tr>
<tr>
<td>Kenyon et al</td>
<td>Botswana</td>
<td>Antituberculosis fixed-dose combination</td>
<td>13</td>
<td>NR</td>
<td>TLC, LC, and UV</td>
<td>NR</td>
<td>4/13 (31%)</td>
<td>Low and high API%</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: API, active pharmaceutical ingredient; NR, not reported; HPLC, high-performance liquid chromatography; TLC, thin-layer chromatography; NIR, near-infrared spectroscopy; UV, ultraviolet spectroscopy; LC-MS, liquid chromatography-mass spectrometry; NMR, nuclear magnetic resonance spectroscopy; IR, infrared spectroscopy; TB, tuberculosis.
medicines such as warfarin and levothyroxine.\textsuperscript{67} Two studies from Eastern Europe found some problems regarding low API% and dissolution failures when a limited number of antibiotics were analyzed in Estonia and Russia.\textsuperscript{83,84} No studies could be identified that addressed medicine quality problems in the Australian continent.

### Neglected noncommunicable medicines in SSFFC surveys

Most of the studies in this review were found to explore medicines used to treat infectious diseases such as malaria and tuberculosis. Medicines used to treat noninfectious diseases, also known as noncommunicable disease (NCD) medicines or chronic disease medicines, were only found in a few studies that presented some medicine quality problems.\textsuperscript{31,32,47,58–60,67,77,80} However, on a global scale, NCDs and their medicines must not be ignored. The WHO estimates that NCDs kill more than 36 million people each year, of which 29 million deaths (80%) occur in low- and middle-income countries.\textsuperscript{97} The currently available literature on medicine quality does not reflect the wider use of NCDs and their medicines globally, including in lower-income countries. This issue needs to be addressed rapidly, as recent evidence from Pakistan reported the death of more than 100 people after the administration of the antianginal medicine isosorbide mononitrate contaminated with large amounts of pyrimethamine.\textsuperscript{98,99} Elsewhere, the US Food and Drug Administration recently issued warnings regarding counterfeit cancer medicines.\textsuperscript{100,101} Furthermore, evidence of counterfeiting involving NCD medicines such as diabetes treatments were found in illicit or lifestyle drugs, which may have significant implications for the public health and could result in death.\textsuperscript{17,102,103} Therefore, it is recommended that we extend the attention of future medicine quality surveys globally beyond infectious diseases medicines and on to NCD medicines (and widely available treatments of diabetes and cardiovascular diseases in particular), in addition to cancer treatments and narrow therapeutics index medicines, as they could have severe health implications for the affected population.

### Type of analysis used in SSFFC surveys

All studies included in this review performed chemical analysis for the identification and/or quantification of the API available in selected samples, in accordance with our methodological approach. High-performance liquid chromatography and TLC were the most widely used chemical analytical techniques available in the selected articles, possibly because of their wide acceptance in the academic field and their application in many pharmacopeial references. It is suggested that this would be a logical and possibly important consideration for future scholars interested in conducting medicine quality surveys to ensure the acceptance of their findings within the academic field.

Physical analysis tests were performed to complement chemical analysis in approximately two-thirds of the selected studies, particularly disintegration and dissolution tests for solid dosage forms. This can be attributed to the availability of specific physical tests in different pharmacopoeias in addition to the use of physical information about the medicinal product to predict the bioavailability of medicines.\textsuperscript{2,45,88} However, such physical analysis tests could only be used as a bioavailability indicator and cannot substitute lengthy and expensive bioavailability studies.\textsuperscript{89,93} Moreover, it is important to note that performing physical analysis only on medicinal samples can be considered inadequate if the objective of the study was to determine medicine quality issues, as it cannot be determined whether the correct API and its quantity are present in medicine samples, as specified in the WHO definition of substandard and counterfeit medicines.\textsuperscript{3–5}

Package inspection is another popular type of medicine analysis that was also found in nearly two-thirds of the medicine quality surveys in this review. On the basis of primary and secondary package information, the majority of reports seek obvious spelling errors, suspicious holograms compared with known genuine samples, and basic label misinformation such as medicine name, dosage, manufacturer details, expiry date, and lot number (Tables 1, 2, and 4). The WHO definition of counterfeit medicines highlights packaging information significance and could have influenced the wide use of package information among medicine quality surveys.\textsuperscript{7} Furthermore, packaging information of medicines has been a valuable mode of analysis in the relevant literature and has revealed many counterfeit medicines that have passed chemical identification tests.\textsuperscript{34,41,43} A tool kit developed by the World Health Professions Alliance and the International Pharmaceutical Federation for visual inspection of medicines can be used for a systematic package inspection by health care professionals and scholars both in practice and in future investigative projects.\textsuperscript{104}

A less common level of analysis available in the literature is the authentication of medicine source via contact with the medicine manufacturer and local or international health authorities. We have identified only ten research articles that attempted to authenticate the source of the medicine samples.\textsuperscript{12,33,35,40,46,56,62,80,86,88} Perhaps researchers...
may not guarantee adequate responses to their queries from other parties, as some have suggested.\textsuperscript{33,35} It could also be possible that authenticating the source may not be within the scope of a particular medicine quality survey, as it could be only focused on substandard medicines issue.\textsuperscript{51} Nevertheless, the WHO definition of counterfeit medicines clearly describes the deliberate and fraudulent misrepresentation of the medicine source as a characteristic of a counterfeit medicine.\textsuperscript{3} Moreover, according to the Pharmaceutical Security Institute, counterfeit medicines are currently increasing in terms of reported incidences worldwide and can no longer be ignored.\textsuperscript{105} We recognize that obtaining authentication confirmation of medicine sources could be difficult in studies collecting samples from street markets; however, this task could be less complex when samples are collected from pharmacies or hospitals, as official records and documentation of medicines are expected to exist. Furthermore, according to the limited studies that reported authentication analysis in this review, many counterfeit cases were found by confirmation from manufacturers or health authorities of a nonauthentic batch of medicines, even if samples contained the correct API when chemically analyzed.\textsuperscript{35,46,86}

Overall, there were very few research articles that performed all four levels of analysis: chemical, physical, package inspection, and authentication of source.\textsuperscript{33,35,40,62,86,88} Future medicine quality surveys are advised to consider performing all four types of analysis for a more holistic approach, and equally, to address the possibility of finding either counterfeit or substandard medicines during an investigation. Further, it was noted that none of the medicine quality surveys examined patient information leaflets within medicinal packages to check for accuracy and up-to-date information made available to patients. Some studies, particularly in the Middle East, have found disagreement between patient information leaflets in some medicine samples when compared with national formularies.\textsuperscript{106,107} Therefore, the addition of patient information leaflets to examination of medicine samples in medicine quality survey studies is open for debate among the scientific community.

Prevalence of SSFFC

Our data suggest that reports of substandard medicines are more widely available in the literature, particularly medicines with incorrect API% and failure of dissolution/disintegration tests, than counterfeit medicine reports (Tables 1–4). These findings are in line with previous reports that suggested that substandard medicines are more prevalent than counterfeits and require more global attention.\textsuperscript{23,24} This phenomenon might be attributed to poor manufacturing practices or extreme weather conditions in some countries, accompanied by inadequate storage conditions.\textsuperscript{3,5,8} However, because the majority of cited articles in this review did not conduct authentication processes via contact with manufacturers and/or health authorities, as previously mentioned, medicine counterfeiting remains a possibility that has not been largely explored. Hence, considering the available data, it cannot be determined whether substandard medicines are indeed more prevalent than counterfeit medicines at this time. Future medicine quality researchers are therefore encouraged to remain vigilant about counterfeiting possibility and conduct all types of analysis including chemical, physical, package inspection, and authentication efforts to determine the type of medicine quality problem more accurately.

Limitations of this review

This systematic review is not without limitations. Articles conducting chemical analysis were a prerequisite for inclusion in this review. We focused only on prospective field quality surveys and excluded reporting of any studies with retrospective or previously seized SSFFC medicines in the literature. Studies proposing novel chemical or physical analytical techniques and methods are typically conducted on previously seized samples of SSFFC medicines, and therefore would not be covered within this review. Our search strategy has limited our findings to the search terms used and the databases searched. We did not search for articles on the Internet in an attempt to preserve the systematic nature of our study. The Internet source of medicines was beyond the scope of our review. Relevant articles from the bibliographical list of available studies were only included on some occasions and cannot be considered exhaustive.

The included articles were not assessed for the quality of their methodology, which was found to vary considerably among the selected articles. The primary author was the only individual who performed the identification, selection, and inclusion of articles in this review. No attempt was made to calculate prevalence rates of SSFFC medicines or test for statistical significance, as it would have resulted in the exclusion of most articles from this review, as most reported studies used convenience sampling and/or with limited sample size.\textsuperscript{11}

Strengths of this review

This review has several strengths. To our knowledge, it is only the second systematic review on the subject of
SSFFC medicines. Evidence of SSFFC medicines in terms of nature and type of analysis were discussed. This information would most likely aid government agencies and health care authorities and scientists interested in the medicine quality issues in developing or improving current policies and practices. It was the intention of this review to help interested parties identify and describe SSFFC medicine problems with up-to-date scientific evidence. Further, this review highlighted neglected medicine types and neglected geographical location in terms of scientific research addressing SSFFC medicines. This could invite more research projects addressing these neglected medicines and geographical locations to improve current knowledge on the issue and maintain patient safety. Moreover, this review has identified the limited scientific research, conducting field quality surveys on SSFFC medicines, using all four levels of analysis, in an attempt to encourage future researchers to explore all possibilities when conducting a medicine quality survey in any settings.

Conclusion
The problem of SSFFC medicines is evident worldwide. Potential harm to patients’ health requires global collaboration exceeding the status quo. Limited research addressing SSFFCC medicines was noted in several parts of the world, including the Middle East, North Africa, and Australia. Similarly, more research is required to address SSFFC medicines from noncommunicable medicine classes, including narrow therapeutic index and chronic medicines, as current scientific knowledge regarding these medicines remains limited despite their popularity and media reports of the existence of SSFFC medicine problems in such therapeutic classes. Furthermore, the current focus of published research on chemical and physical analysis of medicine samples could overlook the possibility of counterfeiting if additional steps of analysis were performed, including package inspection and authentication of source via contact with manufacturers and health authorities. Future medicine quality surveys are encouraged to perform all four levels of analysis to explore all possibilities of substandard and counterfeit medicines that may be present in their selected sample of medicines. Such an approach would be beneficial in determining the type and prevalence rate of medicine quality problems in any setting and could consequently determine the most appropriate strategies to combat their threats.

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