Performance of LED fluorescence microscopy for the detection of acid-fast bacilli from respiratory samples in peripheral laboratories in Argentina

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Introduction: Light-emitting diode fluorescence microscopy (LED-FM) has been endorsed by the World Health Organization (WHO) for tuberculosis diagnosis, but its accuracy in HIV-infected patients remains controversial, and only some few studies have explored procedural factors that may affect its performance.

Objective: To evaluate the performance of LED-FM for tuberculosis diagnosis in patients with and without HIV infection using a newer, less expensive LED lamp.

Materials and methods: We compared the performance of LED-FM and Ziehl-Neelsen (ZN) microscopy on respiratory specimen smears from tuberculosis (TB) suspects and patients on treatment examined by different technicians blinded for HIV-status and for the result of the comparative test. We analyzed the effect of concentrating specimens prior to microscopy using different examination schemes and user-appraisal of the LED device.

Results: Of the 6,968 diagnostic specimens collected, 869 (12.5%) had positive Mycobacterium tuberculosis cultures. LED-FM was 11.4% more sensitive than ZN (p<0.01). Among HIV-positive TB patients, sensitivity differences between LED-FM and ZN (20.6%) doubled the figure obtained in HIV-negative patients or in those with unknown HIV status (9.3%). After stratifying by direct and concentrated slides, the superiority of LED-FM remained. High specificity values were obtained both with LED-FM (99.9%) and ZN (99.9%). The second reading of a sample of slides showed a significantly higher positive detection yield using 200x magnification (49.4 %) than 400x magnification (33.8%) (p<0.05). The LED-device had a very good acceptance among the technicians.

Conclusion: LED-FM better performance compared with ZN in HIV-infected patients and user-appraisal support the rapid roll-out of LED-FM. Screening at 200x magnification was essential to achieve LED-FM increased sensitivity.

Key words: Tuberculosis; fluorescence; microscopy; diagnosis; sputum; acid-fast bacilli.

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As of 2011, the World Health Organization has recommended the use of LED fluorescence microscopy (FM) as an alternative of Ziehl-Neelsen (ZN) in a phased manner for a more rapid tuberculosis (TB) diagnosis (1). FM offers many advantages over ZN. It can detect approximately 5-10% more acid-fast bacilli (AFB) positive smears (2) than ZN. The higher contrast of AFB fluorescence allows the screening of slides at a much lower magnification (200X or 400x) than ZN (1,000X), and they can be examined more quickly, which offers potential solutions to the high workload in some laboratories (3). Besides, FM staining quality may be easier to control because the quality of commercial auramine dye is less variable than that of basic fuchsin (4). Furthermore, the staining technique is simpler and the auramine solution is easier to prepare (5). As no immersion objective is needed to observe the bacilli, there is no need to use the immersion oil and the xylene required with ZN, which are both expensive and may damage the objectives through mishandling or due to poor quality (6). However, fluorescent microscopes using mercury-vapor lamps (MVL) are relatively expensive, they have a short life span and require a reliable electricity supply, besides, replacement bulbs may be difficult to obtain (1). These factors have led to an interest in FM using LED. Compared to conventional MVL fluorescence microscopes, LED microscopes are less expensive and have fewer maintenance requirements. The diodes are very durable, do not require warm-up time, and do not contain toxic products. More importantly, they are reported to perform equally well without a darkroom (1).

As HIV-infected patients are considered a population in whom ZN microscopy tends to produce a low yield, interventions to increase the sensitivity of microscopy are needed. Conventional MVL-FM has shown promising results in HIV-infected individuals; two relevant studies have reported that FM was significantly more sensitive for the diagnosis of pulmonary TB than ZN microscopy in this group of patients (7,8). On the other hand, accuracy data for LED-FM in HIV-infected patients are scarce (9-12); furthermore, the results of some studies are conflicting with those obtained with conventional FM, showing a similar sensitivity (10) or lower specificity (9, 11) values for LED-FM compared with ZN smear microscopy. In addition, only few studies have evaluated the possible effect of sputum processing on the performance of LED-FM, as well as the best scheme for smear examination to achieve increased sensitivity (10,12,13). Besides, as available commercial LED systems may have different operational characteristics, to scale up the replacement of ZN by LED-FM more research is needed to evaluate the different LED systems recently introduced in the market. In low and middle income countries, local production and technology transfer is a good strategy to increase access to medical devices. The TK-LED microscope lamp (Tolket S.R.L.), produced by a technology company based in Buenos Aires, Argentina, uses the latest high power LED illumination technology, it is compatible with most leading microscope brands and can be adapted to any of them on requirement.
Argentina has a well-structured TB laboratory network which performs approximately 150,000 AFB smear microscopy examinations every year. The technical quality and agreement of ZN smear microscopy has been satisfactory over the last years, but operational and epidemiological analyses have shown that there is a need of increasing the number of sputa studied by microscopy in order to prevent diagnostic delay (14,15). The replacement of light microscopy by FM would be one of the immediate options to cope with expected increases in workload, especially in high-burden settings.

This study evaluated the performance of LED-FM for the detection of AFB from respiratory samples of patients with and without HIV infection using a newer, less expensive national LED lamp mounted on an Olympus CX31 microscope, taking culture as the reference standard. We also evaluated different LED-FM technical requirements under field conditions.

Materials and methods

Participating laboratories and technicians

In 2012, seven public laboratories were included in a multicenter project to assess the feasibility of using LED-FM in Argentina. Two of these laboratories had previous experience in conventional FM, and one of them was designated as coordinating center responsible for training, monitoring, slide rechecking and data management. The project was developed in two phases: (i) The technicians training phase, and (ii) the LED-FM evaluation study itself. The first phase was held during 2012, whereas the assessment study was accomplished during 2013 and 2014.

Laboratories were selected according to their workload, HIV prevalence among TB patients and interest in the project. Besides, to be eligible the site had to have acceptable performance in ZN blinded smear rechecking process during the last three years. A ZN acceptable performance laboratory was defined as that with less than 5% error rate and no high false positive or high false negative errors.

The technicians involved in the project attended a three-day course followed by a two-month on-site training in which each technician examined slides from the same specimen stained by ZN and FM in unblinded manner in order to acquire confidence to recognize the bacillus. During this training phase, technicians proficiency was evaluated using a monthly testing panel (composed of nine negative and nine positive smears with different positivity degrees), and a monthly random blinded rechecking. Trainees were evaluated only if they showed an acceptable performance defined as follows: 1) At the most, one low false negative (LFN) error, i.e., a scanty (1-19 AFB/1 length) positive smear misread as negative, without any high false negative results (a 1+ to 3+ positive smear misread as negative) or false positive errors (FP), i.e., a negative smear misread as positive in the last proficiency testing panel, and 2) at the most two minor errors (no more than one LFP, i.e., a negative smear misread as a scanty one, 1-19 AFB /1 length) without any major error (high false positive or high false negative) in the rechecking process.

Clinical specimens included in the evaluation study and smear microscopy

This study was conducted using specimens submitted routinely to the participating laboratories for mycobacterial culture. We included a total of 6,968 diagnostic and 2,419 follow-up respiratory specimens (comprising sputum, bronchialveolar lavage, bronchial wash and lung aspirates).

Direct or concentrated smears (16) were prepared in duplicate following the method routinely used in each participating laboratory. Hot ZN technique (0.3% carbolfuchsine and 0.1% methylene blue) (17) or 0.1% auramine O (counterstained with 0.5% potassium permanganate) for LED-FM (18) were used for staining each of these slides. ZN slides were examined with the bright field microscopy used routinely in each laboratory at 1,000X magnification, whereas FM slides were examined with Olympus CX31 microscopes with a TK-LED illumination (Tolket S.R.L., Buenos Aires, Argentina), using 200X magnification for screening and 400X magnification for confirming and quantifying the slides. The laboratory technicians were also advised to read doubtful slides using higher magnification (1,000X) for confirmation. LED-FM grading of smears was done according to WHO/IUTLD guidelines (18), whereas ZN grading was made according to national guidelines (17) (table 1). After examination, all smears were kept in a dark room at ambient temperature (23°-27°C). After smear preparation, all specimens were decontaminated, concentrated by centrifugation and the deposits were inoculated on solid and/or liquid media according to the technique routinely established in each participating laboratory and following the national tuberculosis guidelines on culture (16). The result of diagnostic samples culture was used as the reference standard. Once contaminated
specimens or those in which a non-tuberculous mycobacterium was isolated were excluded, 6,637 diagnostic and 2,419 follow-up specimens were finally analyzed. HIV status data was extracted from the registers kept in the participating laboratories.

A similar proportion of slides stained by ZN and FM were read by the microscopists unaware of patients’ characteristics and comparative tests results. In one of the laboratories, in which only one technician was responsible for reading both slides, blinding was ensured by labelling FM and ZN slides with different identification numbers; besides, different registers were used to record ZN and FM results.

The external quality assurance protocol as specified by national guidelines for ZN microscopy (17) was in place in all these services. For the purpose of this multicenter evaluation, FM performance was assessed by rechecking all slides at the coordinating center laboratory. Slides were re-stained and re-read in a blinded manner using two different examination schemes: (A) Two lengths at 200X magnification with 400X confirmation, and (B) one or two lengths at 400X. Two distinct LED devices, TK-LED and Primostari-LED (Carl Zeiss, Jena, Germany), were used to re-read these slides in a blinded manner. The decision of enriching our sample with low positive smears (50% of scanty and 50% of negative slides) was made in an effort to target those slides most at risk of being missed by the different reading schemes, in case there would be any difference in the accuracy of both checking diagrams. We chose to evaluate the schemes by using the Zeiss device in conjunction with TK-LED microscope considering the wide acceptance of the Zeiss microscope in global evaluations.

End-user appraisal

A qualitative end-user appraisal survey was conducted among laboratory technicians after approximately three months of experience with LED-FM. The questionnaire included rating of signal-noise ratio, uniformity and intensity of fluorescent illumination, ease of focusing and scale-up suitability.

Statistical analysis

Sensitivity and specificity values were calculated for FM and ZN using culture as the reference standard for diagnosis specimens. Those specimens contaminated in culture were excluded for the analysis as diagnosis could not be made with certainty. We calculated ZN and FM positivity rates for specimens collected from follow-up patients on anti-tuberculosis treatment.

Statistical analysis was performed using the Epidat 3.1, version (PAHO) and Medcalc (MedCalc Software bvba). Results were considered significant at p<0.05. Comparison of sensitivity, specificity and positivity rates in samples from follow-up patients where both methods were applied to the same specimen were done using the McNemar’s test. We also calculated positive and negative predictive values with 95% confidence intervals (CI) for each microscopy test.

Subgroups were set by HIV infection and by concentrating or not specimens before smear preparation; a chi-square test was used to compare sensitivities in subgroups.

To compare semi-quantitative microscopy results with LED-FM and ZN, we calculated CI, and statistical significance was defined as non-overlapping 95% CI.
McNemar’s test was used to compare ZN and LED-FM positivity obtained by different examination schemes or using different LED devices applied to the same sample of smears.

Ethical review was waived as there was no potential risk to participants’ safety, privacy or confidentiality since no formal contact occurred between investigators and participants either directly (interview, questionnaires, etc.) or indirectly (medical records, personal identifiers, etc.). Data on respiratory specimens provided for routine clinical care services were completely anonymized before their inclusion in the study.

Results

LED-FM and ZN performance

Out of the 6,637 diagnostic specimens with microscopy results, culture and species confirmation, 869 (12.5%) were culture-positive for *M. tuberculosis* complex. ZN and LED-FM performance in the seven participating laboratories is shown in table 2. Overall, the sensitivity of LED-FM was 11.4% higher than that of ZN (87.7%; 762/869 vs. 76.3%; 663/869; McNemar’s test: p<0.001), while the levels of specificity obtained with both methods were high and quite similar (99.9% for ZN vs. 5,763/5,768; 99.9% for LED-FM; McNemar’s test: p=1). We found a statistically significant difference in sensitivity between the two methods in all laboratories (McNemar’s test: p<0.05) except for laboratory #5 (McNemar’s test: p=0.22). The overall positive and negative predictive values for *M. tuberculosis* detection by LED-FM were 99.4% (95%CI 98.4-99.8) and 98.2% (95%CI 97.8-98.5), respectively, whereas the corresponding values for ZN were 99.4% (95%CI 98.4-99.8) and 96.6% (95%CI 96.0-97.0), respectively.

In follow-up patients, the positivity rate for LED-FM (14.2%; 344/2,419) was 3.2% higher than for ZN (11.0%; 267/2,419; p<0.001 McNemar’s test).

Diagnostic sensitivity in HIV-infected individuals

Table 3 shows that LED-FM was more sensitive than ZN, irrespective of HIV status (p<0.001 for the comparison of sensitivity values of LED-FM vs. ZN in both HIV positive or HIV negative/unknown patients, McNemar’s test). Overall, both ZN and LED-FM had a lower sensitivity among HIV-infected patients compared with most probably HIV-negative patients (59.4% vs. 80.1% for ZN in HIV-infected patients and HIV-negative patients, respectively, chi-square test: p<0.001; 80.0% vs.

### Table 2. Comparison of ZN and LED-FM performance for TB diagnosis using culture as reference standard

<table>
<thead>
<tr>
<th>Laboratory</th>
<th>TP/C+ Sensitivity</th>
<th>Increase in sensitivity</th>
<th>p value (McNemar’s test)</th>
<th>TN/C- Specificity</th>
<th>p value (McNemar’s test)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory #1</td>
<td>ZN 69/93 74.2</td>
<td>9.7</td>
<td>0.004</td>
<td>820/821 99.9</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 78/93 83.9</td>
<td></td>
<td></td>
<td></td>
<td>820/821 99.9</td>
<td></td>
</tr>
<tr>
<td>Laboratory #2</td>
<td>ZN 125/156 79.5</td>
<td>8.3</td>
<td>0.003</td>
<td>1,452/1,453 99.9</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 137/156 87.8</td>
<td></td>
<td></td>
<td></td>
<td>1,452/1,453 99.9</td>
<td></td>
</tr>
<tr>
<td>Laboratory #3</td>
<td>ZN 163/193 84.5</td>
<td>11.9</td>
<td>&lt;0.001</td>
<td>823/823 100.0</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 186/193 96.4</td>
<td></td>
<td></td>
<td></td>
<td>822/823 99.9</td>
<td></td>
</tr>
<tr>
<td>Laboratory #4</td>
<td>ZN 47/57 82.5</td>
<td>15.8</td>
<td>0.004</td>
<td>850/850 1</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 56/57 98.2</td>
<td></td>
<td></td>
<td></td>
<td>850/850 1</td>
<td></td>
</tr>
<tr>
<td>Laboratory #5</td>
<td>ZN 59/88 67.0</td>
<td>5.7</td>
<td>0.22</td>
<td>381/381 1</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 64/88 72.7</td>
<td></td>
<td></td>
<td></td>
<td>380/381 99.7</td>
<td></td>
</tr>
<tr>
<td>Laboratory #6</td>
<td>ZN 83/108 76.8</td>
<td>10.2</td>
<td>0.01</td>
<td>714/714 1</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 94/108 87.0</td>
<td></td>
<td></td>
<td></td>
<td>714/714 1</td>
<td></td>
</tr>
<tr>
<td>Laboratory #7</td>
<td>ZN 117/174 67.2</td>
<td>17.4</td>
<td>&lt;0.001</td>
<td>724/726 99.7</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 147/174 84.5</td>
<td></td>
<td></td>
<td></td>
<td>725/726 99.9</td>
<td></td>
</tr>
<tr>
<td>Total of specimens</td>
<td>ZN 663/869 76.3</td>
<td>11.4</td>
<td>&lt;0.001</td>
<td>5,764/5,768 99.9</td>
<td>1</td>
</tr>
<tr>
<td>LED-FM 762/869 87.7</td>
<td></td>
<td></td>
<td></td>
<td>5,763/5,768 99.9</td>
<td></td>
</tr>
</tbody>
</table>

TP: True positive; C+: *M. tuberculosis* culture-positive specimens; TN: True negative; C-: Culture-negative specimens; ZN: Ziehl-Neelsen; LED-FM: Light emitting diode fluorescence microscopy
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LED fluorescence microscopy in TB diagnosis

89.4% for LED-FM in HIV-positive patients and HIV-negative patients, respectively, chi-square test: p=0.002) (table 3.)

HIV positive and negative culture-positive specimens were stratified according to the method used for smear preparation. As shown in table 3, both ZN and FM sensitivities seemed slightly lower, though not significantly, in direct smears than in concentrated smears; among HIV patients, ZN direct and concentrated smear sensitivities were 55.3% vs. 60.7%, respectively (chi-square test: p=0.69), whereas values for LED-FM using direct and concentrated smears were 73.7% and 82.0%, respectively (chi-square test: p=0.38). Similarly, among most probably negative HIV-patients, ZN sensitivity was slightly lower using direct smears (79.9%) in comparison with concentrated smears (82.7%) (chi-square test: p=0.76), while LED-FM sensitivities using direct or concentrated smears were very similar (89.3% vs. 90.4%, respectively, chi-square test: p=1.00)

Table 3. Comparison of the ZN and LED-FM sensitivities stratified by HIV in direct and concentrated smears using culture as reference standard

<table>
<thead>
<tr>
<th>Specimens from HIV positive patients</th>
<th>Specimens from negative or unknown HIV status patients</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TP</strong></td>
<td><strong>C+ Sensitivity (%)</strong></td>
</tr>
<tr>
<td>ZN</td>
<td>95</td>
</tr>
<tr>
<td>LED-FM</td>
<td>100</td>
</tr>
<tr>
<td><strong>Direct smears</strong></td>
<td></td>
</tr>
<tr>
<td>ZN</td>
<td>21</td>
</tr>
<tr>
<td>LED-FM</td>
<td>28</td>
</tr>
<tr>
<td><strong>Concentrated smears</strong></td>
<td></td>
</tr>
<tr>
<td>ZN</td>
<td>74</td>
</tr>
<tr>
<td>LED-FM</td>
<td>100</td>
</tr>
</tbody>
</table>

TP: True positive; C+: M. tuberculosis culture-positive specimens; ZN:Ziehl-Neelsen; LED-FM: Light emitting diode fluorescence microscopy

chi square test for the comparison of ZN and LED-FM sensitivities among HIV positive and negative/unknown patients; results presented for all specimens and stratified by treatment of sample before smear preparation (direct/concentrated smears)

ZN positive in direct smears not significantly lower compared with ZN positive in concentrated smears, chi square test: p=0.69

LED-FM positive in direct smears not significantly lower compared with LED-FM positive in concentrated smears, Chi-square test: p=0.38

ZN positive in direct smears not significantly lower compared with ZN positive in concentrated smears, chi square test: p=0.76

LED-FM positive in direct smears not significantly lower compared with LED-FM positive in concentrated smears, chi square test: p=1.00

Comparison of microscopy results with LED-FM and ZN

A total of 227 specimens had discrepant results: 25 slides were positive using ZN and negative using LED-FM. Another 202 slides that were positive in LED-FM were reported negative using ZN (table 4).

When low positive smears (scanty and +) were analyzed, 197 of the 504 (39.1%, 95% CI 34.9-43.4%) detected by LED-FM were negative by ZN, while 25 of the 361 (6.9%, 95% CI 4.7-10.2%) detected by ZN were negative by LED-FM.

Table 4. Comparison of microscopy results with LED-FM and ZN

<table>
<thead>
<tr>
<th>ZN</th>
<th>Negative</th>
<th>Scanty</th>
<th>1+</th>
<th>2+</th>
<th>3+</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>LED-FM</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>7,920</td>
<td>154</td>
<td>43</td>
<td>5</td>
<td>0</td>
<td>8,122</td>
</tr>
<tr>
<td>1+</td>
<td>25</td>
<td>85</td>
<td>175</td>
<td>75</td>
<td>1</td>
<td>361</td>
</tr>
<tr>
<td>2+</td>
<td>0</td>
<td>3</td>
<td>35</td>
<td>130</td>
<td>85</td>
<td>253</td>
</tr>
<tr>
<td>3+</td>
<td>0</td>
<td>2</td>
<td>7</td>
<td>51</td>
<td>260</td>
<td>320</td>
</tr>
<tr>
<td>Total</td>
<td>7,945</td>
<td>244</td>
<td>260</td>
<td>261</td>
<td>346</td>
<td>9,056</td>
</tr>
</tbody>
</table>

ZN: Ziehl-Neelsen

LED-FM: Light emitting diode fluorescence microscopy
Overall, there was an increase in the proportion of positivity in all categories of smear quantification with the use of LED-FM; of the 9,056 specimens included in the study, the number of positives increased from 361 slides (4.0%; CI 3.6-4.4%) with ZN to 504 smears (5.6%; CI 5.1-6.1%) with LED-FM for low positive smears, and from 573 slides (6.3%; CI 5.8%-6.8%) with ZN to 607 smears (6.7%; CI 6.2-7.2%) with LED-FM for high positive smears (++ and +++).

Comparison of examination schemes

There was no difference in smear-positive detection yield between TK-LED and Primostari-LED; (McNemar’s test: p=1) (table 5). On the contrary, the proportion of positivity with both microscopes was significantly higher when using reading scheme A (two length at 200x/ 400x for confirmation) in comparison with examining one or two lengths at 400x magnification (scheme B) (table 5).

End-user appraisal

All the fourteen technicians participating in the study gave positive feedback on TK-LED device signal-noise ratio, uniformity of fluorescent illumination and ease to focus. All believed that the use of LED-FM could improve examination speed and that the TK-LED microscope could be used without a darkroom. All technicians said that LED-FM could be scaled up, and twelve thought that laboratories with a big daily workload should be given priority in the employment of LED-FM. No technical problems were reported during the 36 months of TK-LED device usage both for the study and in routine practice in two laboratories.

Discussion

Our findings further confirm the previously reported better performance of LED-FM over the conventional ZN technique (19,20,21). A significant increased sensitivity of LED-FM over ZN was found in all laboratories except in laboratory #5. Although it is well known that microscopy performance is highly dependent upon factors such as the setting and the population attended in the service (22,23), we identified no factor that could bias the results in favor of ZN performance in laboratory #5. On the other hand, it is unlikely that such result could be explained by a poor proficiency of the technicians working in this service, as they showed an acceptable performance during the training phase and the rechecking process. Furthermore, given the small sample size in this laboratory, the power of the statistical test was about 35%, which suggests that there was not a very high chance of detecting differences between both tests, even though LED-FM showed a higher sensitivity.

Among HIV patients, two relevant studies (7,8) performed with conventional FM reported increased sensitivity over ZN. Nevertheless, although published studies have shown that LED microscopy works as well as conventional FM, both in research and in operational settings (19,24,25), the few studies evaluating LED-FM utility for diagnosing TB in HIV-infected people have shown doubtful results. Two studies (9,11) have reported a similar sensitivity of LED-FM compared to ZN, and one of them, conducted in Indonesia(11), showed that FM was less specific than ZN. Conversely, our results are in line with those performed with conventional FM, showing that among HIV-positive TB patients, the difference in sensitivity between LED-FM and ZN (20.6%) doubled that obtained in the group of TB patients with negative or unknown HIV status (9.3%). However, a recent report (26) showed that LED-FM sensitivity in the HIV-infected population continued to be lower than that obtained in most probably HIV-negative patients, possibly due to the higher occurrence of paucibacillary TB in HIV population and the difficulties in obtaining good quality sputum specimens in advanced cases (27).

As various methods of concentrating sputum based on centrifugation have been shown to increase

**Table 5.** Smear-positive yield in 160 slides read using different reading schemes and two LED devices

<table>
<thead>
<tr>
<th></th>
<th>Number of positive smears among 160 slides (%)</th>
<th>P value (McNemar’s test) (a)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Two lengths at 200x 400X for confirmation (A)</td>
<td>One length at 400X (B1)</td>
</tr>
<tr>
<td>iLED positivity</td>
<td>79 (49.4)</td>
<td>39 (24.4)</td>
</tr>
<tr>
<td>TK-LED positivity</td>
<td>79 (49.4)</td>
<td>38 (23.8)</td>
</tr>
<tr>
<td>p value (McNemar’s test) (b)</td>
<td>1.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>

NA: Not applicable
(a) p values for the comparison of the proportion of positivity using A reading scheme with both B1 and B2 schemes
(b) p values for the comparison of the proportion of positivity obtained by i-LED and that obtained with TK-LED using different reading schemes
diagnostic yield when used prior to microscopy (28), the use of direct or concentrated smears could behave as a confounding factor when evaluating LED-FM performance in different populations.

In agreement with results recently reported by Getachew, et al.(12), when we stratified smears by their direct/concentrated state, the diagnostic superiority of LED-FM over ZN remained equivalent. FM appears to be a more sensitive technique than ZN due to its ability to detect low bacillary load in the sputum, as previously reported (11,21). In accordance with these studies, we found that the use of LED-FM significantly increased the proportion of low positive smears in about 40.0% compared with ZN. A stronger absorbability of mycolic acid by auramine-O than by fuchsine would partially explain the increased sensitivity of LED-FM for AFB. Furthermore, according to results by other researchers (20,29), applying both microscopy techniques to follow-up sputum, expected to have an increased proportion of scanty results compared to diagnostic sputum (30), the positivity rate with LED-FM was significantly higher than with ZN. Other factors that contribute to this higher performance of LED-FM, such as improvement in the proficiency of the laboratory technicians due to additional training in FM and new microscopes, may have played a role to a lesser extent, especially considering that one of the conditions for including a laboratory in this multicenter study was an acceptable performance in the ZN external quality assessment during the last three years.

We found that even LED-FM had a higher sensitivity than ZN, as 25 ZN-positive specimens resulted negative by LED-FM. This may be explained by two phenomena: First, as AFB are not homogeneously distributed in sputum, the portion of sputum used to prepare ZN-stained slide may have contained them, whereas the portion used to prepare the slide stained with auramine may have not, especially in those specimens with scarce bacilli. On the other hand, the uneven distribution of AFB on the slide may also explain this phenomenon; as technicians can only observe a proportion of fields on a slide, some AFB may have not been detected simply because the technicians examined those fields that were randomly AFB-depleted; this can explain why different laboratory professionals examining the same number of fields on a slide may get differing results, as it has been previously reported (31). The larger field area subject to examination using FM compared to the oil immersion fields needed when ZN is used may also explain FM higher sensitivity. The field of view when a 20x objective is used is three to four times larger than that of the 40x objective and 25 times larger than that of the 100x objective used for ZN microscopy (32). Accordingly, we found that, in agreement with Kubica’s study (33) using conventional FM, scanning with 200x magnification enabled a higher positivity yield than examining smears at 400x only, while AFB remained clearly visible with both of the stand alone FM systems used in this study. On the contrary, other authors have reported that using two LED modules mounted on routine microscopes screening smears under 200x magnification did not increase sensitivity compared to screening under 400x magnification (34,35). These authors also reported that with these devices AFB were less clearly visible under 200x compared to 400x magnification, which may explain the differences between their results and ours when LED-FM technology but different devices were evaluated.

The impossibility of comparing TK-LED and Primostari-Led performance was a limitation of our study. To ensure a more precise comparison of positive detection by the two LEDs using different reading schemes, we analyzed a random selection of low positives and the same number of negatives detected with TK-LED. In this small fraction of TK-LED negative slides, no false negatives were detected using the i-LED device. However, given the low proportion of negatives evaluated with both microscopes, it does not seem unreasonable to assume that no TK-LED false-negatives would have been detected if all the negatives had been reread with the i-LED microscope. In these circumstances we were able to compare the detection capacity with the two reading schemes, but not the overall performance of the two LED devices.

Concerns have been voiced regarding the possibility of false positive smears with LED-FM (due to impurities in auramine, food particles and artifacts which produce some fluorescence), and it has been suggested that all scanty and doubtful cases must be confirmed by ZN (36). However, this tends to oversee the increased sensitivity and efficiency gained with the use of LED-FM, which some experts have recommended to discourage (37). The level of specificity of LED-FM in our study was high and similar to that obtained with ZN. Conversely, in their evaluation of LED-FM in HIV-infected patients, Chaidir, et al. (11), reported a significantly lower specificity compared to ZN microscopy, and they pointed out that false positive LED-FM results may have been due to lack of experience or training.
among technicians. Albert, et al. (9), reported a marked difference in the specificity of LED-FM among readers recently trained to evaluate LED-FM performance in HIV-infected patients, which highlights the importance of implementing careful quality assurance measures when a new technology is introduced.

Our study had these additional limitations: (i) it was performed by specimen and not by patient, and while this is consistent with most other studies in this field, we acknowledge that the lack of independence between specimens from the same patient may have led to overestimate accuracy, and (ii) HIV testing was not done systematically as part of the study; according to laboratory records, no HIV status had been registered for a significant number of patients, which underpowered the result analysis by HIV status.

Regarding feasibility aspects, we confirmed the wide acceptance of LED-FM by technicians and the possibility of using it without a dark room. We also found that new national instruments worked well during the first three years. These are all very significant factors in scaling-up LED-FM in national TB programs with limited resources.

To conclude, our findings confirmed LED-FM better performance compared to ZN in HIV-infected patients as previously recorded for conventional FM. Furthermore, user appraisal supported LED-FM rapid roll-out, especially when the burden of HIV-infected population is high. The use of 20x objective for screening was essential to achieve increased sensitivity. WHO has recently endorsed the implementation of Xpert MTB/RIF assay as the first initial TB screening in HIV patients (38), however, microscopy is still needed for treatment monitoring. In their comparison of LED-FM and Xpert MTB/RIF assay in HIV patients, Álvarez-Uria, et al. (39), reported that although the molecular assay showed an increase in positive results in comparison with LED-FM when performing both tests in the same specimen, this increase was only modest when analyzing two specimens by LED-FM. Besides, the cost of the equipment, the annual maintenance and the consumables are considerably higher for the Xpert MTB/RIF assay, and this could hinder its use in some resource-limited settings. The 2015 TB Report (40) granted that LED-FM adoption remained low; in 2014, this technology was reported to be present in only 2% of microscopy centers in the Region of the Americas. In this sense, we hope our findings are useful to increase the implementation of this technology. Particularly in Argentina, the operational and technical evidences we provided will be a valuable input in the preparation of a national strategic plan to expand the use of LED-FM.

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Conflicts of interest

The authors have declared that no conflicts of interest exist.

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References

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