A Strategy for Screening and Confirmation of HTLV-1/2 Infections in Low-Endemic Areas

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Serological tests have been widely used for detecting human T-cell lymphotropic virus type 1/2 (HTLV-1/2) antibodies in the endemic areas, but their performance in low-risk populations is rarely reported. The aim of this study was to evaluate the performance of four HTLV-1/2 screening assays and to discuss a strategy for diagnosis of HTLV-1/2 infection in a non-endemic area. At the present study, 1546 specimens repeatedly reactive (RR) by one screening ELISA were collected from blood centers/banks from January 2016 to April 2019. Avioq-ELISA, Murex-ELISA, Roche-ECLIA and Fujirebio-CLIA were independently performed on each plasma sample and compared to WB and LIA confirmatory tests. Positive or indeterminate specimens with blood available were quantified by qPCR. The results showed that 48 samples were finally confirmed as HTLV-1 positive, 13 were HTLV positive, 151 were indeterminate, and 387 were negative. All the WB-positive samples were also LIA-positive. Roche-ECLIA showed the highest sensitivity that was able to detect 91.8% positives and combined with the Murex-ELISA would significantly increase the positive detection rate (98.4%). In addition, LIA yield more indeterminate and HTLV-untyped results than WB (152 vs. 27), but was able to resolve infection status of some individuals with an indeterminate WB. Besides, 3 WB indeterminate and 1 LIA-untyped samples were confirmed as HTLV-1 positive by qPCR. Based on these findings, we put forward a proper test strategy for HTLV-1/2 diagnosis in low-prevalence areas. If possible, the Roche-ECLIA with the highest sensitivity is suggested as a second screening assay in primary labs. If not, all RR specimens are recommended to be firstly retested by Roche-ECLIA and Murex-ELISA in the reference lab. Secondly, samples reactive to any one of the two tests were quantified by qPCR, and then the NAT-negatives were furtherly submitted to LIA for confirmation. Thereby, the cost can be reduced and the diagnostic accuracy would be improved.

Keywords: T-cell lymphotropic virus type 1/2 (HTLV-1/2), enzyme-linked immunosorbent assay (ELISA), chemiluminescence immunoassay (CLIA), electrochemiluminescence immunoassay (ECLIA), western blot, line immunoassay
INTRODUCTION

Human T-lymphotropic virus (HTLV) was the first RNA retrovirus to be associated with cancer (Poiesz et al., 1980). To date, four HTLV related viruses (types 1 to 4) have been discovered, but only HTLV-1 and HTLV-2 have been convincingly linked to human diseases (Poiesz et al., 1980; Kalyanaraman et al., 1982; Calattini et al., 2005; Wolfe et al., 2005). HTLV-1 is the etiologic agent of Adult T-cell leukemia (ATL) and HTLV-associated myelopathy/tropical spastic paraparesis (HAM/TSP) (Verdonck et al., 2007). HTLV is mainly spread via mother-to-child transmission, sexual contact, and through contaminated needles shared by drug users. It also can be transmitted through the transfusion of infected blood components and tainted liver, kidney, or lung transplants. To mitigate these risks, mandatory screening of blood supplies for HTLV-1/2 was implemented in the mid-1980s in most developed and several developing countries (Williams et al., 1988; Maeda, 1989). However, because most regions of China have long been considered non-endemic for HTLV, limited screening has been implemented.

Currently, initial diagnosis of HTLV-1/2 infection is mainly based on detecting specific antibodies in plasma or serum using enzyme-linked immunosorbent assay (ELISA) and CLIA or ECLIA. Several commercial kits based on recombinant and/or synthetic peptide antigens alone or in combination with viral lysates have also been adapted for large-scale screening of HTLV-1/2 antibodies. However, the kits fail to differentiate between HTLV-1 and HTLV-2 infections because the two types share a high homology between them (Andersson et al., 1999; da Silva Brito et al., 2018). Besides, the high false-positive rate of these commercial assays, especially in low-seroprevalence populations, is a major problem. Therefore, confirmatory assays with high specificity are required for samples exhibiting a signal repeatedly greater than or equal to the threshold value of the screening assays.

Western blot is most frequently used to confirm the presence of anti-HTLV-1/2 antibodies. The FDA-licensed MP Diagnostics HTLV Blot 2.4 uses a combination of recombinant HTLV-1/2 proteins and HTLV-1 viral lysate to improve sensitivity. This assay also uses HTLV type-specific recombinant envelope protein (gp46-1 and gp46-2) to discriminate viral types (Varma et al., 1995). However, a large number of WB-indeterminate and WB-untypeable results are commonly found in several population groups and especially among low-risk blood donors (Mahieux et al., 2000; Ando et al., 2004; Yao et al., 2006). To address this shortcoming, another serologic confirmatory test with improved sensitivity, INNO-LIA HTLV, has been developed to confirm and differentiate HTLV-1 and HTLV-2 infections (Zrein et al., 1998). This assay appeared useful in reducing the numbers of inconclusive WB results (Zrein et al., 1998).

Molecular assay is another method used for HTLV confirmation. Real-time PCR or quantitative PCR (qPCR) have been used to determine the real status of HTLV infections and quantify the proviral load (PVL). However, these molecular assays have had low sensitivity in special population groups, such as individuals infected with HIV or HTLV-2 (Montanheiro et al., 2008; Campos et al., 2017).

Although several commercially available HTLV-1/2 diagnostic assays have been used worldwide, their performance in endemic and non-endemic areas varies greatly. Moreover, few studies have been conducted to compare the performance of LIA and WB in low-prevalence populations. Therefore, the objective of this study was to evaluate several FDA- or CE-licensed HTLV-1/2 assays to diagnose the specimens that were RR to one ELISA, and to discuss strategies for the diagnosis of HTLV-1/2 infection in a non-endemic area.

MATERIALS AND METHODS

Samples and Study Design

All specimens were collected from eligible donors from blood banks/centers in 26 provinces in China between January 2016 and April 2019. The RR samples were delivered at 2–8°C or frozen to the NCCL for confirmation. The RR samples were also HBsAg, anti-HCV, anti-HIV, and anti-TP negative by both two screening ELISAs in the enrolled blood centers. According to the confirmatory process (Figure 1), all RR samples were simultaneously subjected to four screening assays in the NCCL. Samples reactive to any one of the four assays were further confirmed by a LIA (INNO-LIA HTLV I/II score, Fujirebio, Japan) and WB (MP HTLV Blot 2.4, MP Biomedicals, Singapore).

Samples with limited plasma volumes that were not enough for all assays were excluded from our study. Enrolled samples with a positive result either by WB or LIA were classified as positive. Positive or indeterminate specimens with whole blood available were further quantified by qPCR in our lab. In addition, blood donations with an indeterminate result were recollected from the same donor, and retested by the same detection process.

Screening and Evaluated Assays

Three different kinds of commercially available ELISA assays [Wantai HTLV-1/2 antibody ELISA kits (Wantai BioPharm, China), Murex HTLV I+II (Diasorin S.p.A., United Kingdom) and Foresight HTLV-1/2 antibody ELISA kits (Acon Biotech, China)] were used for screening HTLV1/2 antibodies at all participating blood centers/banks in China. Reactive specimens were retested by the same assay and any one of two-round retests found to be reactive were defined as RR specimen.

Further detection was done in the NCCL using Elescsys HTLV-I/II (Roche Diagnostics, Germany), Lumipulse G HTLV-I/II (Fujirebio, Japan) and two kinds of ELISA assays (Murex HTLV I+II (Diasorin S.p.A., United Kingdom) and Avioq HTLV-I/II Microelisa System (Avioq, Durham, NC, United State). The key features of the evaluated assays are summarized in Table 1.

Abbreviations: CLIA, chemiluminescence immunoassay; ECLIA, electrochemiluminescence immunoassay; ELISA, enzyme-linked immunosorbent assay; HTLV-1/2, human T-cell lymphotropic virus type 1/2; LIA, line immunoassay; NCCL, National Center for Clinical Laboratories; PPV, positive predictive value; RR, repeatedly reactive; WB, western blot.
FIGURE 1 | The testing algorithms of HTLV-1/2 in our study. WB, western blot; ELISA, enzyme-linked immunosorbent assays; CLIA, chemiluminescence immunoassays; ECLIA, electrochemiluminescence immunoassays; RR, repeatedly reactive. +: positive/reactive; –: negative/non-reactive; +/-: indeterminate.

TABLE 1 | Technical features of evaluated assays and confirmatory assays used in the study.

| Assay name          | Manufacturer      | Assay type                                          | Detection antigens                              | Testing time |
|---------------------|-------------------|----------------------------------------------------|-------------------------------------------------|--------------|
| Elecsys HTLV-I/II   | Roche diagnostics | One-step double-antigen sandwich chemiluminescent immunoassay | Viral recombinant antigens gp21 and p24          | 18 min       |
| Lumipulse G HTLV-I/II | Fujirebio        | Two-step sandwich chemiluminescent immunoassay    | Recombinant antigens p19 I/II, p24 I/II, gp46 I/II, gp21 I | 4 min        |
| Avioq HTLV-I/II Microelisa System | Avioq | ELISA                                              | Viral antigens (purified viral lysate) and recombinant HTLV-1 p21E antigen | 150 min      |
| Murex HTLV-I/II     | Diasorin          | ELISA                                              | HTLV-1 and HTLV-2 antigens                       | 90 min       |
| INNO-LIA HTLV I/II score | Fujirebio | A line immunoassay                                | Recombinant antigens p19 I/II, p24 I/II, gp46 I/II, gp21 I/II | 18 h         |
| MP HTLV Blot 2.4    | MP Biomedicals    | Western Blot assay                                 | Recombinant HTLV-1/2 antigens and HTLV-1 viral lysate | 150 min      |

ELISA, enzyme-linked immunosorbent assay.

**Confirmatory Tests**

In the LIA, the strips contained recombinant proteins (rp19 I/II, rp24 I/II, and rgp21 I/II) for confirmation and synthetic peptides (gp46 I, p19 I, and gp46 II) for differentiation of HTLV-1 and HTLV-2 antibodies. Samples were considered HTLV positive if they were reactive with at least two confirmation bands, including...
The positive samples were typed as HTLV-1 if the intensities of gp46I and p19I bands were equivalent to or higher than that of the gp46II band. HTLV-2 positivity was defined if the gp46II band was more intense than the gp46I and p19I bands and if no discriminate bands appeared, an untypable HTLV result was denoted. A specimen was classified as indeterminate if it were reactive with one band (gp21I/II) or two bands (except rgp21I/II). Samples were considered negative if a single rp24I/II, rp19I/II, gp46I, or gp46II band or no bands appeared.

WB is most frequently used as a supplemental assay for confirming and differentiating the presence of antibodies against HTLV. Results of MP HTLV Blot 2.4 were interpreted according to the stringent criteria provided by the manufacturer. Briefly, HTLV-1 seropositivity was defined as reactivity to gag (p19 with or without p24) and two env (GD21 and rgp46-I). HTLV-2 seropositivity was defined as reactivity to gag (p24 with or without p19) and two env (GD21 and rgp46-II). Samples that were reactive to both gag (p19 and p24) and env (GD21) were defined as HTLV seropositive but were considered untypable. Any other patterns of specific bands that did not meet the above criteria were considered as indeterminate. Absence of bands or appearance of non-specific band patterns were interpreted as negative.

Quantitative PCR (qPCR)
DNA was extracted from whole blood of each specimen using the Tiangen Magnetic Blood Genomic DNA Kit (Tiangen Biotech, China). HTLV-1 proviral DNA was detected and quantified by qPCR using a set of primer pairs and a TaqMan probe targeting the HTLV-1 pol region. The RPPH1 gene was also amplified simultaneously as an internal control. The primers and probes are shown in Supplementary Table S1. Standard curves were generated using recombinant plasmid DNA with HTLV-1 pol and RPPH1 sequences. Normalized HTLV-1 PVL was calculated using the formula: (HTLV-1 DNA average copy number/RPPH1 average copy number) × 2 × 100 leukocytes and expressed as the number of HTLV-1 copies per 100 PBMCs. The detection limit for the HTLV-1 pol region was estimated to be 2.5 copies per 10,000 host leukocytes with a 95% hit-rate.

Statistical Analysis
The PPVs were calculated by SPSS version 21.0 (IBM, Corp., Armonk, NY, United States). Descriptive statistics are presented as geometric means + standard deviations. Results were analyzed with t-test. p < 0.05 was considered statistically significant.

RESULTS
HTLV Confirmation and Typing
A total of 1546 RR samples with enough volumes were included in the final sample count in our study. Of the 1546 samples, 555 showed discordant results and 991 showed consistent results across the four assays. Of the 991 consistent samples, 44 were reactive to all four assays while the rest were negative in all assays. Finally, 599 samples that were reactive to at least one assay were confirmed by LIA and WB. Of these samples, 73.29% (439/599) showed consistent results in the two confirmatory tests, including 44 HTLV-1 positive samples, 8 indeterminate samples and 387 negative samples (Table 2). As samples were defined as positive if any confirmatory test were positive, 48 samples were finally identified as HTLV-1 positive, 13 as HTLV positive, 151 as indeterminate and 387 as negative. In addition, 41 blood samples were also tested by qPCR, which showed that 6 HTLV-1 positive samples and 1 HTLV-untyped samples were NAT-positive and 30 indeterminate samples were NAT-negative. The results and validation algorithm are shown in Figure 1.

As shown in the Table 2, 61 samples were interpreted as positive using the LIA as a reference. The positive detection rates for the four screening assays ranged from 72.1 to 91.8% (Table 3 and Supplementary Table S2). A combination of any two of the four assays increased the positive detection rate. We found that the combination of Roche-ECLIA and Murex-ELISA detected 98.4% positives, which was higher than other combinations. However, only 44 samples were classified as positive when WB was used as a standard and the positive detection rates were increased to 93.2, 97.2, 100, and 100% for Avioq-ELISA, Murex-ELISA, Roche-ECLIA, and Fujirebio-CLIA, respectively (Supplementary Table S2).

Correlation Between the PPVs and the Reactivity Index of the Evaluated Commercial Assays
In each assay, positive samples confirmed by LIA and WB displayed a significantly higher reactivity index than indeterminate (p < 0.0001) and negative samples (p < 0.0001) (Figure 2). Moreover, the difference in reactivity indexes between indeterminate and negative samples as detected by the evaluated assays was also statistically significant (p < 0.05), indicating that reactivity indexes may correlate with the confirmatory results.

To determine the relationship between PPVs and reactivity indexes on four evaluated assays, we analyzed the results of the assays at different cut-off reactivity indexes for the four assays (Table 4). We found that when the cut-off values were 1.0, the PPV for Avioq-ELISA was 91.7%, but for Murex-ELISA, Roche-ECLIA and Fujirebio-CLIA the PPVs were only 29.5, 21.0, and 23.7%, respectively. PPVs above 95% were
TABLE 3 | Details of the discrepant results by LIA and WB among the finally confirmed HTLV positive samples.

| Sample no. | ELISA Avioq S/CO Murex S/CO Roche S/CO Fujirebio COI | CLIA 427.9 3.15 | LIA Bands pattern | WB 2.4 Results | WB 2.4 PVL | qPCR Results |
|------------|-----------------|------------------|------------------|----------------|-----------|--------------|
| 1626       | 1.787           | 3.315            | 50.0             | p19 I/II, gp24 I/II, gp46 I/II, gp21 I/II, gp46-I | HTLV-1 | 0.339 HTLV-1 |
| 1705       | 1.166           | 3.297            | 50.0             | p19 I/II, gp24 I/II, gp46 I/II, gp21 I/II, gp46-I | HTLV-1 | 11.397 HTLV-1 |
| 359        | 5.323           | 4.099            | 50.0             | p19 I/II, gp24 I/II, gp46 I/II, gp21 I/II, gp46-I | HTLV-1 | / NT |
| 1739       | 1.146           | 3.49             | 50.0             | gp46 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 525        | 0.14            | 0.43             | 3.1              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 495        | 0.19            | 0.40             | 8.4              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 1441       | 0.34            | 0.55             | 5.6              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 1747       | 0.074           | 2.02             | 0.6              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Ind / NT |
| 53         | 0.142           | 2.212            | 1.8              | p24 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 1412       | 0.17            | 2.59             | 0.4              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 816        | 0.24            | 1.86             | 0.3              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 1295       | 0.15            | 1.02             | 0.9              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 489        | 0.16            | 0.35             | 5.18             | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 647        | 0.13            | 0.45             | 4.15             | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 1285       | 0.29            | 0.36             | 1.32             | p24 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 1461       | 0.32            | 0.38             | 1.47             | p24 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |
| 457        | 0.190           | 0.780            | 3.1              | p19 I/II, gp21 I/II, gp46-I | HTLV-1 | / Neg / NT |

observed when cut-off values were 1.5 for Avioq-ELISA, 10.0 for Murex-ELISA, 29.0 for Roche-ECLIA and 8.8 for Fujirebio-CLIA. PPVs were 100% when the cut-off ratios for Avioq-ELISA, Murex-ELISA, Roche-ECLIA and Fujirebio-CLIA were 2.5, 11.0, 67.2, and 28.0, respectively.

FIGURE 2 | S/CO or COI values distribution among the finally confirmed positive, indeterminate and negative samples. Ind, indeterminate; Pos, positive; Neg, negative.
Discrepancies Between INNO-LIA and WB in the Finally Confirmed Positive Samples

Seventeen samples that were finally defined as HTLV-1 or HTLV positive showed discrepant results between INNO-LIA and WB (Tables 2, 3). Of these 17 samples, 3 were LIA HTLV-1 positive and 14 were LIA HTLV positive but untypable. Two out of the 3 LIA HTLV-1 positives were also NAT-positive but were WB-indeterminate while the other one was WB-negative. One out of the 14 LIA-untyped samples was discriminated as HTLV-1 by WB, 4 were confirmed as WB-indeterminate and 9 were WB-negative. It is noteworthy that 1 LIA-untypable but WB-indeterminate sample was also NAT-positive.

After combining the results confirmed by LIA, WB and qPCR, 5 samples were classified as HTLV-1 positive and 12 samples were HTLV positive but untypable (Table 3). The average S/CO or COI values of the untyped HTLV infections for Murex-ELISA, Roche-ECLIA and Fujirebio-CLIA were 1.03, 5.49, and 2.09, respectively. All these untyped samples did not respond to Avioq-ELISA and their positive rates for Roche-ECLIA, Murex-ELISA and Fujirebio-CLIA were 7/12, 5/12 and 5/12, respectively, indicating that these specimens may easily be missed by some screening assays.

Characterization of Indeterminate Results

Of the 151 samples which were finally interpreted as indeterminate, 130 were LIA ± /WB-, 15 were LIA-/WB ± and 6 were indeterminate by both tests (Tables 2, 5). Most of the LIA-indeterminate samples showed a gp21 I/II pattern while only a few displayed a p19 I/II plus p24 I/II pattern. Thirty LIA-indeterminate samples were available for qPCR detection, but none were positive. In the WB-indeterminate samples, a single gp46 I/II band was the most frequently observed pattern, followed by a single GD21 and a single p19 band. Of these indeterminate samples, 68.7% were Roche-ECLIA positive, 54.3% were Fujirebio-CLIA positive, 43.7% were Murex-ELISA positive and only 0.4% were Avioq-ELISA positive.

Thirteen indeterminate samples were recollected from the same donors no less than 8 weeks after last donation and retested by all assays (Supplementary Table S3). Seroconversion occurred in 2 individuals with LIA ±/WB- results. Both samples converted from the single gp21 I/II band pattern to the gp21 I/II plus p19 I/II band pattern but were still WB-negative. In addition, 4 out of the 13 indeterminate samples, including 2 LIA ±/WB- samples and 2 LIA-/WB ± samples, converted to negative. Six recollected samples were also retested by qPCR, but all of them were NAT-negative, including the seroconverted LIA-HTLV positive sample.

### DISCUSSION

Currently, screening for HTLV-1/2 antibodies in healthy blood donors is not done routinely in China. A systematic review revealed that HTLV-1/2 prevalence in the coastal areas of China, and especially in Fujian province, is higher, compared to the low average HTLV-1/2 prevalence in the mainland (Chen et al., 2019). However, it is notable that in some provinces close to the high-endemic areas of China, including Hunan, Jiangsu and Guangxi, the prevalence is relatively high, probably due to large-scale population migration in these regions in recent years (Chen et al., 2019). Accordingly, HTLV-1/2 may have spread from the coastal areas to the mainland, resulting in increased opportunities for virus transmission and infection. Therefore, it is reasonable to expect HTLV-1/2 detection to become necessary in China. Consequently, appropriate strategies for screening and confirming HTLV-1/2 infection in China should be developed urgently.

In our study, almost one-third samples displayed discordant results among the four evaluated assays, which may due to the differences in the combination, source and quantity of HTLV-1/2 antigens used by different manufacturers. None of the evaluated assays could detect all the true HTLV positives when using LIA as a standard. In contrast, the Roche-ECLIA was a relatively more sensitive test that was reactive to 91.8% confirmed positive samples. This assay was especially sensitive to the untyped positives (7/12, 58.33%). Moreover, Roche-ECLIA could detect more indeterminate samples than the other three assays, indicating it was a sensitive method for screening HTLV-1/2 antibodies. From the perspective of blood banks/centers, sending RR samples to a reference lab is costly and time-consuming. Therefore, we propose that, where possible, such samples could be retested by the Roche-ECLIA and the positives subjected to subsequent confirmatory tests. Our data suggest that this would reduce the need for confirmation by about two thirds.

The present study indicated that most untypable HTLV infections had relatively low S/CO or COI values or were misdiagnosed by some screening tests, indicating a high risk of missed detections when only one screening assay is used in the reference labs. In our study, the combination of Roche-ECLIA and Murex-ELISA performed relatively well, detecting 98.4% true positives when LIA was the standard. We detected 100% positive samples when Murex-ELISA was combined with Roche-ECLIA and Fujirebio-CLIA but this strategy would be costly, time-consuming and labor intensive for reference labs.

HTLV-1-associated diseases develop in only a small proportion of infected individuals and slowly progress to
disease relative to the average life span of humans. Consequently, after balancing screening/conformational costs and the threat of HTLV infections to public health especially in the non-endemic area, we recommend the use of a combination of Roche-ECLIA and Murex-ELISA in reference centers to retest initially reactive blood samples in low-risk areas. Since Roche-ECLIA is more sensitive than Murex-ELISA, it is best to test RR samples with Roche-ECLIA first and then the negatives by Murex-ELISA. Therefore, the number of test would be reduced by about 12% compared with test simultaneously.

Our results also indicate that higher S/CO or COI values were associated with higher diagnostic reliability. As demonstrated in this study, samples with S/CO values over 2.5, 11.0, 67.2 for Avioq-Murex, Murex-ELISA, and Roche-ECLIA, respectively, or COI value over 28.0 for Fujirebio-CLIA, were rarely negative. Hence, samples with values above these thresholds can be interpreted as HTLV-1/2 positive, decreasing the demand and cost of confirmatory tests. Accordingly, if Murex-ELISA and Roche-ECLIA were combined for sample retesting in reference centers, the number of samples needing confirmation would reduce by 14.7%. Nevertheless, molecular assays or other supplementary tests would be necessary to determine PVL or HTLV infection types.

WB-indeterminate and/or untypable variants remain a serious problem worldwide. It has been found that the frequency of WB-indeterminate samples reactive to HTLV-1 ELISA varies among different cohorts, but is much higher in low-risk blood donors (Martins et al., 2010; Abrams et al., 2011). In our study, only 4.5% (27/599) of WB-indeterminate samples and no WB untypable samples were identified. The low sensitivity of WB and the special specimens collected from blood donors in the HTLV-1 infected area may account for these results. Of the WB-indeterminate samples, 22.2% (6/27) were confirmed positive by LIA and 11.1% (3/27) were further confirmed HTLV-1 positive by qPCR. Although most of the WB-indeterminate, LIA-positive samples (4/6) were LIA untypable, the results indicated that LIA was more sensitive than WB and, in our view, should be the first choice for confirmation.

Indeterminate WB patterns indicating true or false virus infection have been reported previously (Martins et al., 2010). In our study, rgp46-II alone was the most frequently observed pattern, which was remarkably different from the pattern observed in blood donors or general population from other regions (Hayes et al., 1991; Mauclère et al., 1997; Lu and Chen, 2003; Filippone et al., 2012), indicating that indeterminate WB patterns can vary among populations. Martins et al. (2010) reported that the presence of p19 plus p24 seemed to indicate true reactivity and p24 alone indicated false reactivity. In our study, the p19 band was observed in 5/6 WB ± /LIA+ samples, 3 of which also exhibited another unique band (p24 or rgp46-I or GD21). Therefore, we suspected that the presence of p19 plus a unique GAG or ENV band indicated true HTLV infection and should be validated by LIA or molecular methods. In high-risk or endemic areas, indeterminate WB patterns may represent seroconverters (Jacob et al., 2008; Martins et al., 2010).

In this study, seroindeterminate blood donors were usually recalled for recollection. However, only three WB-indeterminate samples were returned, two of which showed faint rgp46-II bands and were reassigned as negative. We estimated that a proportion of the WB-indeterminate samples were false reactions, but larger follow-up studies to test this notion should be performed.

Although LIA reduced the number of WB-indeterminate samples, the assay yielded more indeterminate (22.7%, 136/599) and untypable (2.3%, 14/599) results than WB. Most of the samples expressed a single gp21 I/II pattern (97.1%, 132/136), the largest of which had faint intensity (55.1%, 75/136). We propose several reasons for these inconclusive results, including low HTLV proviral loads, mutations in the provirus (defective particles), seroconversion period or cross-reactivity with other antigens or viruses. In our study, 30 LIA-indeterminate and 5 LIA-untyped samples were tested by qPCR, but only one

### TABLE 5 | The reactivity patterns of the indeterminate samples.

| Ind. pattern | INNO-LIA reactivity pattern | WB reactivity pattern | No. | qPCR results (positive no./ total no.) | No. of samples positive by the following assay/ total no. of samples tested: |
|--------------|-----------------------------|----------------------|-----|---------------------------------------|--------------------------------------------------------------------------------|
|              |                             |                      |     |                                       | Avioq | Murex | Roche | Fujirebio |
| INNO-LIA±/WB−| gp21 I/II                   | /                    | 127 | 0/30                                  | 4/130 (0.3%) | 54/130 (41.53%) | 93/130 (71.5%) | 72/130 (55.38%) |
|              | p19 I/II,p24 I/II           | /                    | 3   | NT                                    | 2/15 (12.3%) | 8/15 (53.3%) | 9/15 (60.0%) | 5/15 (33.3%)   |
| INNO-LIA−/WB±| /                           | rgp46-II GD21        | 5   | NT                                    | 0/6 (0.00%) | 4/6 (66.67%) | 4/6 (66.67%) | 5/6 (83.3%)    |
|              | /                           | p19                 | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
|              | /                           | p24                 | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
|              | /                           | p19,p26             | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
|              | /                           | rgp46-I             | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
|              | /                           | p24, rgp46-I        | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
| INNO-LIA±/WB±| gp21 I/II                   | GD21                 | 3   | NT                                    | 0/6 (0.00%) | 4/6 (66.67%) | 4/6 (66.67%) | 5/6 (83.3%)    |
|              | gp21 I/II                   | rgp46-I             | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
|              | gp21 I/II                   | rgp46-II            | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
|              | p19 I/II,p24 I/II           | p19,p26             | 1   |                                       | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)  | 1/1 (100%)    |
| Total        |                             |                     | 151 | 0/30                                  | 6/151 (0.4%)| 66/151 (43.7%)| 103/151 (68.7%)| 82/151 (54.3%) |

LIA, line immunoassay; WB, western blot; ±, indeterminate; −, negative; NT, not tested.
LIA-untyped sample was confirmed HTLV-1 positive with a PVL of 0.009 copies/100 PMBC, indicating that samples with very low PVL may be hard to identify. Manns et al. (1991) demonstrated that antibodies to rgp21 antigen appeared earlier than antibodies to p24 and p19, and the presence of the single rgp21 band may indicate an early HTLV infection. In our follow-up results, 2 out of 11 recollected samples with strong gp21 I/II intensity converted to HTLV-positive, suggesting a proportion of the LIA-indeterminate samples were in the seroconversion phase and needed to be followed-up. However, 2 samples with faint gp21 I/II intensity were reassigned as negative, revealing that some of the indeterminate samples may have resulted from non-specific reactivity or cross-reactivity with other antigens. Interestingly, samples with strong rgp21 I/II intensity displayed significantly higher S/CO or COI values than samples with faint intensity (Supplementary Figure S1). Taken together, the correlation between higher reactivity indexes and higher diagnostic reliability and the observation that the 2 LIA-indeterminate samples with strong rgp21 I/II intensity retested HTLV-positive, support the view that LIA-indeterminate samples with strong intensities are more likely to still be in the seroconversion period, and should be paid serious attention.

Although the qPCR method used in our study did not detect all the specimens, our results indicated that qPCR was useful for resolving inconclusive confirmatory tests. Due to the large numbers of uncertain results and the high cost of WB and LIA assays, we propose employing qPCR first, followed by WB or LIA to test any qPCR-negatives. Other studies also recommend this strategy, which has been shown to reduce costs and improve the accuracy of HTLV-1/2 diagnosis (Costa et al., 2011; da Silva Brito et al., 2018).

Without a prophylactic vaccine and with limited available treatment options for HTLV-related diseases, recommendations for the deferral and reentry of blood donors are necessary, especially in China where HTLV-1/2 screening is underdeveloped. Based on our study, we suggest that donors with a final positive interpretation by a confirmatory test should be deferred permanently but donors returning a negative result should be reentered. Additionally, donors with indeterminate results should be followed-up, and recollected specimens should be confirmed through the same supplemental test.

**CONCLUSION**

In this study, we compared the performance of four commercially available kits for screening anti-HTLV-1/2 antibodies and used WB and LIA as confirmatory tests. We also proposed a new testing algorithm for screening and confirming HTLV-1/2 in a low prevalence region (Figure 3). We recommend Roche-ECLIA as a second screening test in blood centers/banks to rule out large numbers of false-positives. Where local equipment is lacking, all RR specimens would need to be confirmed in reference labs. To minimize the risk of misdiagnosing true positives, samples that are reactive to Murex-ELISA or Roche-ECLIA should be further confirmed. Taking the cost into consideration, we suggest that these should be quantified using qPCR first, and then the negatives should be subject to LIA but not WB to improve the diagnostic accuracy.
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

ETHICS STATEMENT

This study was approved by the institutional review boards in every participating blood bank. Written informed consent are routinely completed at the time of donation from blood donors.

AUTHOR CONTRIBUTIONS

HJ, LC, and LW designed the study and wrote the manuscript. HJ, LC, XJ, HS, and FG performed the experiments. HJ, LC, YY, and LW analyzed the data. All authors participated in critical revision of the manuscript and approved the final version.

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SUPPLEMENTARY MATERIAL

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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