Survey on common reference intervals for general chemistry analytes in Hong Kong

Toby CH Chan, Chloe M Mak *, Sammy PL Chen, MT Leung, HN Cheung, Daniel CW Leung, HK Lee, Eleanor C Koo, YC Lo

ABSTRACT

Introduction: Reference intervals (RIs) are essential tool for proper interpretation of results. There is a global trend towards implementing common RIs to avoid confusion and enhance patient management across different laboratories. However, local practices with respect to RIs lack harmonisation.

Methods: We have conducted the first local survey regarding RIs for 14 general chemistry analytes in 10 chemical pathology laboratories that employ four different analytical platforms (Abbott Architect, Beckman Coulter AU, Roche Cobas, and Siemens Dimension EXL). Analytical bias was assessed by an inter-laboratory results comparison of external quality assurance programmes.

Results: Sufficient inter-laboratory and inter-platform agreement regarding the 10 analytes (albumin, alanine aminotransferase, aspartate aminotransferase, chloride, gamma-glutamyl transferase, phosphate, potassium, sodium, total protein, and urea) were demonstrated. However, the RIs were heterogeneous across all laboratories, with percentage differences of the upper RI value of up to 47% for aspartate aminotransferase (absolute difference of 16 U/L), 29% for urea (1.8 mmol/L), and 18% for potassium (0.8 mmol/L). The percentage difference between lower RI values was up to 24% for urea (0.6 mmol/L), 22% for phosphate (0.16 mmol/L), and 8% for total protein (5 g/L). The coefficients of variation of the upper RI values of potassium and sodium were 1.2 times and 1.0 times of their corresponding between-subject biological variation, respectively, representing unnecessary variations that are overlooked and unchecked in current practice.

Conclusions: We recommend the use of common RIs for general chemistry analytes in Hong Kong to prevent interpreter confusion, improve electronic data transfer, and unite laboratory practice. This is the first local study on this topic, and our data can lay the groundwork for increasing harmonisation of RIs across more laboratory tests.

New knowledge added by this study

• Reference intervals (RIs) of general chemistry analytes are highly variable.
• Ten analytes (albumin, alanine aminotransferase, aspartate aminotransferase, chloride, gamma-glutamyl transferase, phosphate, potassium, sodium, total protein, and urea) show satisfactory inter-laboratory and inter-platform agreement.
• Implementation of common RIs is feasible.

Implications for clinical practice or policy

• We recommend the use of common RIs in Hong Kong for general chemistry analytes to reduce redundant variation across laboratories.
• This is the first local study on this topic, and our data can lay the groundwork for increasing harmonisation of RIs across more laboratory tests.
引言：参考区间是正确解释结果的重要工具。实施通用参考区间是全球趋势，可避免因诠释引起的混淆并强化不同化验室的患者管理。然而，参考区间间的使用在本地实践中仍缺乏一致性。

方法：我们以10间病理学理化化验室的14种一般化验分析物进行首个有关参考区间的研究。这些化验室采用4种不同的分析平台（Abbott Architect、Beckman Coulter AU、Roche Cobas和Siemens Dimension EXL），通过外部质量保证计划（EQAP）的化验室结果比较评估分析偏差。

结果：其中10种分析物（白蛋白、谷丙转氨酶ALT、天冬氨酸氨基转移酶AST、氮化物、γ-谷氨酰转移酶γ-GT、磷酸盐、钾、钠、总蛋白和尿素）在不同化验室和平台之间有差别的参考区间。在不同化验室之间，AST参考区间上限的百分比差异可达47%（绝对差异16 U/L），尿素为29%（1.8 mmol/L），钾为18%（0.8 mmol/L）。参考区间范围下限的百分比差异可达24%（0.6 mmol/L），磷酸盐为22%（0.16 mmol/L），总蛋白质为8%（5g/L）。钾和钠参考区间上限值的变异系数分别是它们相比较的60%和1.0倍。突显出化验室间参考区间差异的必要性。

结论：我们建议在香港为一般化验分析物订立通用参考区间。这可避免因参考区间不同而引起的解释混乱，改善电子数据传输并统一化验结果。

Introduction

Reference intervals (RIs) are an indispensable tool for clinical decision making in the interpretation of numerical pathology results. Simple yet elegant comparisons with reference subjects empower the interpreter with objective judgements and aid clinicians in diagnosis, treatment, monitoring, prognostication, and screening.1

Reference intervals are commonly defined as limiting values, usually upper and lower limits, between which a prespecified percentage (usually 95%) of results would fall.2,3 In daily practice, for most tests, there exists some degree of laboratory-specific bias related to differences in pre-analytical and analytical factors, such as the choices of analytical platform, methodology, and reagent. Therefore, it is desirable for laboratories to provide sets of laboratory-specific RIs following Clinical and Laboratory Standards Institute guideline C28-A3c. A laboratory may establish a new set of RIs by conducting an RI study with at least 120 reference individuals from each subgroup stratified by sex, age, and other parameters as appropriate.4 Conducting an RI study is challenging, as enormous efforts of human and financial resources are needed. As the list of analytes is long, it is almost impossible for every laboratory to repeat an RI study to accommodate future changes in methodology or analytical platforms.4,5 Alternatively, a laboratory may adopt the RIs established by other sources such as manufacturers or the literature and validate them with at least 20 reference individuals’ results. An additional option is for the laboratory to transfer previously established RIs according to mathematical formulas to account for differences in analytical factors.6 These methods ensure that each laboratory provides a set of clinically meaningful intervals to clinicians, aiding their management.

Therefore, for the same analyte, it is not uncommon to see different RIs across laboratories. This inter-laboratory coefficient of variation was reported by Ceriotti et al7 to be as high as 15% to 20% for the RIs of urea and creatinine. This could be reasonable for hormonal tests that are not optimally standardised, as demonstrated by the marked variations in RIs for thyroid hormones between four analytical platforms shown by a recent study in the UK.8 For analytes that are generally well standardised across platforms, such as plasma electrolytes, one would expect results generated by different laboratories to be comparable. Logically, with insignificant methodological bias, the RIs should be same for the specified homogenous population.

In 2007, the UK Pathology Harmony Group showed that laboratories were using different sets of RIs with no sound scientific basis despite using the same analytical platform and reagents.6,7 The same problem was later also revealed by a survey on RIs in Australasia.9 The differences in RIs were concluded to be unnecessary and would have created unneeded confusion during interpretation, which might lead to inappropriate investigations or treatments.9,10 Common RIs were offered as a solution to unite laboratory practices.4

In Hong Kong, we have observed a general trend of variation in RIs that resembles those in the UK and Australasia, with various RIs adopted for most tests, including general chemistry laboratory tests. Hence, we conducted the first local study to explore the situation with a territory-wide survey on RIs. The aim was to scientifically review the analytical variation of general chemistry laboratory tests between local laboratories and to examine the evidence for such variations.

Methods

Fourteen blood general chemistry analytes were included in this study, namely albumin, alanine aminotransferase (ALT), aspartate aminotransferase (AST), alkaline phosphatase (ALP), total bilirubin, calcium, chloride, creatinine, gamma-glutamyl transferase (GGT), phosphate, potassium, sodium, total protein, and urea. We conducted a territory-
wide survey involving 10 chemical pathology laboratories. All laboratories provided routine services to assess the 14 analytes, except for AST, chloride, and GGT, which were not evaluated in three laboratories. The instruments were Abbott Architect (labs 1-3), Beckman Coulter AU (labs 4-5), Roche Cobas (labs 6-9), and Siemens Dimension EXL (lab 10). Table 1 summarises the analytical platforms and methodologies.

The laboratories participated in the Condensed General Chemistry Programme provided by the Royal College of Pathologists of Australasia Quality Assurance Programs. In each cycle of the external quality assurance programme (EQAP), identical sets of QAP materials were analysed by each individual laboratory for the aforementioned blood general chemistry analytes using their own analytical platform. The use of QAP materials, which were commutable samples with the same properties as routinely analysed clinical samples, minimises the matrix effect. In routine clinical practice, EQAP safeguards laboratory performance by comparison with peers and reference methods. In the present study, we retrospectively review these readily available EQAP data from local laboratories for bias assessments. The participants provided their responses by email to the following items: (1) historical EQAP results of six cycles (105-11, 105-12, 105-15, 105-16, 106-03, and 106-04); (2) adult RIs in use for clinical service, and (3) analytical specification of assays.

Laboratory performance bias was assessed by percentage differences of EQAP results. Percentage difference was defined as the laboratory result minus the target value divided by the target value. The feasibility of applying common RIs among the laboratories was determined by the degree of agreement between the percentage differences and the corresponding allowable limits of performance. Data analyses were performed using Microsoft Excel 2016.

Results

Figure 1 shows that half of the 14 analytes showed agreement across all laboratories. The inter-laboratory differences are within the corresponding target allowable limit of error (ALE) for AST (-3% to +9%; target ALE ±12%), chloride (-1% to +2%; ±3%), phosphate (-1% to 4%; ±8%), potassium (-2% to 3%; ±5%), sodium (-1% to 2%; ±2%). Three other analytes (albumin, ALT, and GGT) also showed agreement.

### TABLE 1. Summary of analytical platform and methodology

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Lab 1</th>
<th>Lab 2</th>
<th>Lab 3</th>
<th>Lab 4</th>
<th>Lab 5</th>
<th>Lab 6</th>
<th>Lab 7</th>
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<td>Beckman Coulter AU</td>
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<td>AMP buffer (other rate</td>
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Abbreviations: ALP = alkaline phosphatase; ALT = alanine aminotransferase; AMP = 2-amino-2-methyl-1-propanol; AST = aspartate aminotransferase; BCG = bromocresol green; BCP = bromocresol purple; Cresol = cresolphthalein; GGT = gamma-glutamyl transferase; GluCANA = L-γ-glutamyl-3-carboxy-4-nitroanilide; IFCC = International Federation of Clinical Chemistry; IMT = integrated multisensory technology; ISE = ion selective electrode; NM-BAPTA = 5-nitro-5’-methyl-BAPTA; P5P = pyridoxine
across nine laboratories with the Abbott, Beckman, and Roche platforms, except Siemens which was only used by one laboratory. Figure 2 shows the inter-laboratory comparison of RIs for the 14 general chemistry analytes. Laboratories using the same platform generally adopted the same RIs, except for one laboratory using the Roche platform.

Notably, for the seven analytes mentioned above that showed agreement within the target ALE, the RIs differed substantially across the 10 laboratories. Particularly, the upper RI limit ranged from 34 to 50 U/L (coefficient of variation [CV]: 11%): in male samples and 30 to 40 U/L (9%) in female samples in AST; 107 to 109 mmol/L (0.9%) in chloride; 1.39 to 1.52 mmol/L (2.7%) in phosphate; 4.4 to 5.2 mmol/L (6.7%) in potassium; 144 to 148 mmol/L (0.7%) in sodium; 79 to 87 g/L (2.2%) in total protein; and 6.3 to 8.1 mmol/L (8.1%) in urea. The lower RIs ranged from 98 to 102 mmol/L (1.7%) in chloride; 0.72 to 0.88 mmol/L (6.2%) in phosphate; 3.4 to 3.6 mmol/L (2.6%) in potassium; 136 to 137 mmol/L (0.2%) in sodium; 63 to 68 g/L (2.2%) in total protein; and 2.5 to 3.1 mmol/L (7.4%) in urea.

The remaining analytes (albumin, ALT, ALP, calcium, creatinine, GGT, and total bilirubin) demonstrated substantial platform-specific bias exceeding the target ALE. High bias exceeding the ALE was observed for ALT (+12% to +20%; target ALE ±12%) and GGT (+11% to +14%; ±12%), with negative bias exceeding the ALE in albumin (-5.3% to -7.1%; ±6%), ALT (-11.4% to -15.3%; ±12%), and calcium (-5.6% to -7.1%; ±4%) present on the Siemens platform. Negative bias exceeding the ALE in ALT (-12.2% to -14.8%; ±12%) was also detected on the Roche platform. For calcium, negative bias exceeding the ALE (-4% to -6%; ±4%) was also detected at one laboratory using the Beckman platform. For creatinine, all laboratories were in agreement about concentrations ranging from 152 to 349 µmol/L. However, significant negative bias (-13% to -22%; ±12%) was observed for creatinine levels at the target value of 67 µmol/L on the Abbott, Siemens, and Roche instruments. For total bilirubin, half of the laboratories showed agreement within the ALE, while the remaining laboratories had significant negative bias (-14% to 17%; ±12%).

The investigated laboratories used different RIs despite employing the same analytical platforms, methods and reagents, for 11 out of the 14 analytes.
among those using the Abbott platform (labs 1-3), 11 out of 14 of analytes among those using Roche platforms (labs 6-9), and three out of the 14 of analytes among those using the Beckman platforms (labs 4-5).

Sex-specific RIs were not consistently provided for eight analytes (ALP, ALT, AST, phosphate, potassium, total bilirubin, total protein, and urea). For instance, sex-specific RIs were not provided by two laboratories for ALP, two for ALT, two for AST, five for potassium, five for urea, seven for total protein, eight for phosphate, and nine for total bilirubin.

**Discussion**

Reference intervals are provided by laboratories as interpretative tools to aid clinical decision making. Theoretically, RIs could be affected by patient factors (eg, sex, age, ethnicity, biological variability), pre-analytical and analytical factors (eg, choice of method, reagents, platform, calibration), and statistical methodology. Therefore, for the same population, the RIs used for a test are inevitably influenced by the bias of the laboratory assays. In other words, RIs should theoretically be the same if the above-listed factors do not introduce significant bias.

In local practice, 10 analytes surveyed demonstrated sufficient agreement within the ALE between different analytical platforms across laboratories (Fig 1: AST, chloride, phosphate, potassium, sodium, total protein, and urea for all four platforms; albumin, ALT and GGT for Abbott, Beckman, and Roche platforms) [Fig 1]. These results confirmed the previous findings of bias assessment by the Australasian Association of Clinical Biochemists, which concluded that chloride, phosphate, potassium, sodium, total protein, and urea measurements showed sufficient similarity across analytical platforms and laboratories and that common RIs could be adopted. The same study found method-specific bias in AST levels averaging +22% for assays using pyridoxal-5-phosphate as an activator compared with those not using pyridoxal-5-phosphate. Our results showed a lesser degree of pyridoxal-5-phosphate–related bias (+7%), so this issue would not prevent the use of common RIs in the local scenario.

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**FIG 2. Inter-laboratory reference intervals of the 14 analytes among the four analytical platforms**

Abbreviations: Abb = Abbott Architect; ALP = alkaline phosphatase; ALT = alanine aminotransferase; AST = aspartate aminotransferase; Bec = Beckman Coulter AU; GGT = gamma-glutamyl transferase; RI = reference interval; Roc = Roche Cobas; Sie = Siemens Dimension EXL.

* The Y-axis refers to the upper and lower RI limits, with square and cross symbols representing those of male and female samples, respectively. The X-axis lists the laboratories in ascending nominal order (ie, labs 1-10) from left to right. Laboratories employing the same analytical platform are highlighted by brackets.
For analytes with demonstrated agreement across platforms and laboratories, the RIs are theoretically expected to be the same if obtained from the same group of reference (ie, ‘healthy’) individuals. In actual practice, for the seven analytes mentioned above, all of the adult RIs varied across laboratories, with the CV of the upper and lower limits of the RIs up to 11% and 7.4%, respectively. The inter-laboratory percentage differences of upper RI limits were up to 47% for AST (absolute difference: 16 U/L), 29% for urea (1.8 mmol/L), and 18% for potassium (0.8 mmol/L), and those of the lower RI limits were up to 24% for urea (0.6 mmol/L), 22% for phosphate (0.16 mmol/L), and 8% for total protein (5 g/L). We can compare the CVs of these analytes’ RIs against the corresponding between-subject biological variation (CV-G) values published by Ricos et al. The CV of the upper RI limits of potassium and sodium were 1.2 and 1.0 times those of CV-G, respectively while that of the lower RI limits of sodium and phosphate were 1.1 and 0.6 times those of CV-G, respectively. These RI variations generate significant additional bias during interpretation, which is often overlooked and unchecked. Furthermore, laboratories were using different RIs despite using the same analytical platforms and methodologies for these analytes. For example, among users of the Abbott platform, the potassium RIs of labs 1 and 2 were 3.6 to 5.2 mmol/L for samples of both sexes, while that of lab 3 was 3.5 to 4.5 mmol/L for male and 3.4 to 4.4 mmol/L for female samples. These variations were unnecessary, as supported by the sufficient agreement across analytical platforms and laboratories. Application of different RIs in various circumstances could lead to confusion among interpreters and hinder data management in the era of electronic health records. Similar trends of unexplained RI variations were previously observed in the UK for sodium, potassium, and other analytes, and this eventually lead to the Pathology Harmony group’s recommendation of harmonised RIs. At present, local laboratories often spend substantial human resources on decisions and maintenance regarding the appropriate RIs for large numbers of analytes. The use of common RIs for these seven analytes would unite local laboratory practices, facilitate electronic communications between laboratory information and electronic patient record systems, and streamline the maintenance of RIs.

For creatinine, low bias was noted for seven laboratories using the Jaffe methods, but this tendency spared the laboratories that used the enzymatic method on the Beckman platform. This bias was likely related to the higher variability of the Jaffe method at low creatinine concentrations, which has been reported to be up to 30% on some platforms. While the remarkably good analytical agreement shown for the remaining five higher concentrations of creatinine support the use of common RIs, this should be cautiously reviewed, as the lowest concentration of creatinine (67 µmol/L) is very close to the lower RI limit. Further study of bias may be warranted for creatinine.

Substantial bias exceeding the ALE was demonstrated for the remaining six analytes, with high bias for ALT and GGT and low bias for albumin, ALP, and calcium on the Siemens platform; low bias for ALP on the Roche platform; low bias for calcium at one laboratory using the Beckman platform; and low bias for total bilirubin in labs 1 to 3, 7, and 8. Positive bias averaging 8% for albumin was observed for laboratories using the bromocresol green method compared with the bromocresol purple method, a pattern similar to the findings of Koerbin et al. Bias in ALT measurement could be attributed to differences in assay design, with an average of +7% bias shown for the assay using pyridoxal-5-phosphate over the assay that does not use it. Bias for calcium and total bilirubin could be related to methodological differences between platforms, while bias for ALP and GGT were likely to be specific to the analytical platform. While the feasibility of local common RIs for these six analytes was not confirmed by this study, our findings indicate that common RIs could still be considered for albumin, ALT, and GGT in laboratories using the Abbott, Beckman, and Roche platforms, which all laboratories except one use.

Variable adoptions of sex-specific RIs were another key finding of the survey. Heterogeneous and inconsistent practices of sex partitioning for RIs were noted in eight analytes (ALP, ALT, AST, phosphate, potassium, total bilirubin, total protein, and urea). Moreover, sex-specific RIs were sometimes different even within the same platform. For example, the upper RI limit of GGT in male samples differed by 11% in laboratories using the Siemens platform; the upper RI limit of ALP differed by 35 U/L among users of the Roche platform, and the upper RI limit of ALP differed by 40 U/L and 7 U/L in male and female samples, respectively, among users of the Abbott platform. Common RIs with united practice of sex partitioning could be the solution to converge these practices.

Historically, heterogeneous and sometimes incomparable results of the same measurands could be obtained with different assays because of suboptimal standardisations in pre-analytical and analytical factors. Laboratory-specific RIs were advocated to compensate and allow for sound interpretations of laboratory results in clinical settings. Realising the need for assay standardisation, an enormous global effort has been taken in the past 60 years to study biological variability, standardise pre-analytical conditions and analytical methods, improve quality control, establish traceability of reference materials and methods, and implement EQA programs for various kinds of assays, led by the International Federation of
Clinical Chemistry (IFCC) and other international/national organisations. Major successes have been realised for a large number of measurands, as listed on the website of the International Consortium for Harmonization of Clinical Laboratory Results.

The concept of common RIs emerged in the early 2000s and has gained huge popularity over the past decade. The theory is simple: if the measured results of different assays are comparable, ie with adequate assay standardisation, the same RIs should be adopted given that the tests are performed on the same reference population. Redundant variations of RIs merely impair interpretation.

Presently, there are two types of common RIs: ‘objective’ and ‘subjective’ ones. Subjective common RIs were generally defined by scientific surveys and expert guidance with the harmonisation approach. Examples include the “agreed Pathology Harmony clinical biochemistry reference intervals for adults” for 15 general chemistry analytes recommended by the UK Pathology Harmony Group in 2011 and the “adult harmonised reference intervals” for 18 general chemistry analytes recommended by the Australasian Association of Clinical Biochemists and endorsed by the Royal College of Pathologists of Australasia in 2016. The two groups have since continued their work on harmonisation of various aspects of pathology in the past decade, with the UK Pathology Harmony Group working on the Pathology Harmony bookmark for tumour markers and requesting guidance for non-specialists, and the Australasian Association of Clinical Biochemists working on harmonisation of paediatric common RIs, serum protein electrophoresis reporting, lipid reporting, management and communication of high-risk lab results, arterial and venous blood gas RIs, and reporting of dynamic endocrine testing for adults and paediatric patients.

Objective common RIs refer to those defined by well-conducted, multicentre reference studies, such as the Nordic Trueness Project, which was conducted with well-standardised pre-analytical and analytical handlings and the use of five control materials. The project involved 102 Nordic routine clinical biochemistry laboratories and more than 2500 carefully selected healthy reference individuals. The Nordic Reference Interval Project RIs for 25 general chemistry analytes were established and published in 2002 and implemented throughout Nordic countries in 2004 with the help of the Scandinavian Society of Clinical Chemistry. Among Asian countries, the Japan Society of Clinical Chemistry has recently published their nationwide common RIs for 40 laboratory tests determined by three multicentre RI studies. Table summarises the common RIs published in different parts of world for the general chemistry analytes surveyed and the common RIs proposed by our study.

In 2017, the IFCC Committee on Reference Intervals and Decision Limits (C-RIDL) published two landmark papers on the results of their global multicentre study on reference values of 25 chemistry analytes in 13 386 healthy adults recruited from 12 countries, including China, with the use of a specially designed serum panel. The study explored the regionality and ethnicity of these reference values globally and provided invaluable information for the possibility of future derivation and transfERENCE of the established RIs through use of the C-RIDL serum panel.

The relatively small number and choice of QAP specimens for retrospective methodological comparisons represent a major limitation of our survey. Artificial materials used in QAP specimens generally gave rise to more variable and method-dependent results due to matrix effects. Despite this, our survey demonstrated that methodological bias would not prevent the use of common RIs for seven general chemistry analytes. For the remaining analytes, we speculate that the degree of methodological bias may be exaggerated by the matrix effect of the QAP, ie, the actual analytical difference is likely to be smaller when tested with a patient sample. Our findings should be verified with a formal prospective bias study with a standardised protocol and the use of another set of blood specimens, preferably unadulterated human samples, with pre-assigned reference values to ensure commutability.

This survey compared the adult RIs of 14 general chemistry analytes among 10 chemical pathology laboratories using four different analytical platforms. Bias assessments and comparisons of RIs revealed that different and variable RIs were provided by the laboratories despite sufficient inter-laboratory and inter-platform agreement regarding the RIs of 10 general chemistry analytes. The use of common RIs was found to be feasible and is recommended for these 10 analytes. Such use would unify and improve local standards of clinical laboratory practice. A well-designed implementation plan for common RIs with support from stakeholders including clinicians, pathologists, and scientists would be vital for the success of such a substantial project. Figure 3 shows our proposed implementation plan for the introduction of common RIs in Hong Kong, modified from the plan suggested by Tate et al for the harmonisation of adult and paediatric RIs in Australasia. Furthermore, the concept of common RIs could be expanded to cover more general chemistry analytes, eg, creatine kinase and magnesium; special chemical tests, eg, therapeutic drug monitoring and hormones; other clinical laboratory specialties, such as haematology and immunology; and paediatric RIs.
TABLE 2. Summary of adult common reference intervals published in United Kingdom, Australasia, Japan, and Nordic countries

<table>
<thead>
<tr>
<th>Analyte (unit)</th>
<th>Sex</th>
<th>United Kingdom$^{21}$</th>
<th>Australasia$^{2,23}$</th>
<th>Japan$^{24}$</th>
<th>Nordic countries$^{28}$</th>
<th>Hong Kong (proposed by this study)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albumin (g/L)</td>
<td>M/F</td>
<td>35-50</td>
<td>-</td>
<td>41-51</td>
<td>36-48 (18-39 years)</td>
<td>36-48 (40-49 years) 34-45 (&lt;70 years) 36-50</td>
</tr>
<tr>
<td>ALP (U/L)</td>
<td>M/F</td>
<td>30-130</td>
<td>30-110</td>
<td>106-322</td>
<td>34-105</td>
<td>-</td>
</tr>
<tr>
<td>ALT (U/L)</td>
<td>M</td>
<td>5-40</td>
<td>10-42</td>
<td>10-70</td>
<td>&lt;54</td>
<td>&lt;40</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>5-35</td>
<td>7-23</td>
<td>10-45</td>
<td>&lt;40</td>
<td>&lt;20</td>
</tr>
<tr>
<td>AST (U/L)</td>
<td>M</td>
<td>5-35</td>
<td>13-30</td>
<td>15-45</td>
<td>&lt;40</td>
<td>&lt;40</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>5-30</td>
<td>15-35</td>
<td>&lt;34</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium (mmol/L)</td>
<td>M/F</td>
<td>-</td>
<td>2.1-2.6</td>
<td>2.18-2.52</td>
<td>2.15-2.51</td>
<td>-</td>
</tr>
<tr>
<td>Chloride (mmol/L)</td>
<td>M/F</td>
<td>95-108</td>
<td>95-110</td>
<td>101-108</td>
<td>-</td>
<td>100-108</td>
</tr>
<tr>
<td>Creatinine (µmol/L)</td>
<td>M</td>
<td>60-110</td>
<td>58-94</td>
<td>60-110</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>45-90</td>
<td>41-70</td>
<td>50-90</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GGT (U/L)</td>
<td>M</td>
<td>5-50</td>
<td>13-64</td>
<td>10-80 (18-39 years)</td>
<td>6-69</td>
<td></td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>5-35</td>
<td>9-32</td>
<td>10-45 (18-39 years)</td>
<td>6-44</td>
<td></td>
</tr>
<tr>
<td>Phosphate (mmol/L)</td>
<td>M/F</td>
<td>0.8-1.5</td>
<td>0.75-1.5</td>
<td>0.9-1.5</td>
<td>0.78-1.44</td>
<td></td>
</tr>
<tr>
<td>Potassium (mmol/L)</td>
<td>M/F</td>
<td>3.5-5.3</td>
<td>3.5-5.2</td>
<td>3.6-4.8</td>
<td>3.5-4.4 (plasma) 3.6-4.6 (serum)</td>
<td>3.5-4.7</td>
</tr>
<tr>
<td>Sodium (mmol/L)</td>
<td>M/F</td>
<td>133-146</td>
<td>135-145</td>
<td>138-145</td>
<td>137-144 (plasma), 137-145 (serum)</td>
<td>136-145</td>
</tr>
<tr>
<td>Total bilirubin (µmol/L)</td>
<td>M/F</td>
<td>&lt;21</td>
<td>1-20</td>
<td>-</td>
<td>5-25</td>
<td></td>
</tr>
<tr>
<td>Total protein (g/L)</td>
<td>M/F</td>
<td>60-80</td>
<td>60-80</td>
<td>66-81</td>
<td>64-79 (plasma), 62-78 (serum)</td>
<td>65-83</td>
</tr>
<tr>
<td>Urea (mmol/L)</td>
<td>M/F</td>
<td>2.5-7.8</td>
<td>-</td>
<td>2.7-7.1</td>
<td>-</td>
<td>2.7-7.5</td>
</tr>
</tbody>
</table>

Abbreviations: ALP = alkaline phosphatase; ALT = alanine aminotransferase; AST = aspartate aminotransferase; GGT = gamma-glutamyl transferase

FIG 3. Proposal for implementing common reference intervals in Hong Kong

Abbreviation: RIs = reference intervals
Author contributions
All authors contributed to the concept or design, drafting of the article, and critical revision for important intellectual content. All authors had full access to the data, contributed to the study, approved the final version for publication, and take responsibility for its accuracy and integrity.

Conflicts of interest
All authors have disclosed no conflicts of interest.

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Ethics approval
The present survey is a retrospective observational review of local laboratory practice and external quality assurance programs designed by the external quality assurance materials used for the data collection in the survey are human body fluid or tissue. The quality assurance program private or sensitive patient data, no collection or analysis of local laboratory practice and external quality assurance programs. The present survey is a retrospective observational review of local laboratory practice and external quality assurance programs designed by the external quality assurance materials used for the data collection in the survey are processed samples designed by the external quality assurance program to mimic the properties of clinical sample. Therefore, ethics approval was not applicable for this study.

References
29. Strowme JH, Rustad P, Steensland H, Theodorsen L, Urdal P. Reference intervals for eight enzymes in blood of adult females and males measured in accordance with the International Federation of Clinical Chemistry reference system at 37 degrees C: part of the Nordic Reference