Quantitative Proteomic Analysis of Plasma Exosomes to Identify the Candidate Biomarker of Imatinib Resistance in Chronic Myeloid Leukemia Patients

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Background: Imatinib (IM), a tyrosine kinase inhibitor (TKI), has markedly improved the survival and life quality of chronic myeloid leukemia (CML) patients. However, the lack of specific biomarkers for IM resistance remains a serious clinical challenge. Recently, growing evidence has suggested that exosome-harbored proteins were involved in tumor drug resistance and could be novel biomarkers for the diagnosis and drug sensitivity prediction of cancer. Therefore, we aimed to investigate the proteomic profile of plasma exosomes derived from CML patients to identify ideal biomarkers for IM resistance.

Methods: We extracted exosomes from pooled plasma samples of 9 imatinib-resistant CML patients and 9 imatinib-sensitive CML patients by ultracentrifugation. Then, we identified the expression levels of exosomal proteins by liquid chromatography-tandem mass spectrometry (LC-MS/MS) based label free quantification. Bioinformatics analyses were used to analyze the proteomic data. Finally, the western blot (WB) and parallel reaction monitoring (PRM) analyses were applied to validate the candidate proteins.

Results: A total of 2812 proteins were identified in plasma exosomes from imatinib-resistant and imatinib-sensitive CML patients, including 279 differentially expressed proteins (DEPs) with restricted criteria (fold change ≥ 1.5 or ≤ 0.667, p<0.05). Compared with imatinib-sensitive CML patients, 151 proteins were up-regulated and 128 proteins were down-regulated. Bioinformatics analyses revealed that the main function of the upregulated proteins was regulation of protein synthesis, while the downregulated proteins were mainly involved in lipid metabolism. The top 20 hub genes were obtained using STRING and Cytoscape, most of which were components of ribosomes. Moreover, we found that RPL13 and RPL14 exhibited exceptional upregulation in imatinib-resistant CML patients, which were further confirmed by PRM and WB.
INTRODUCTION

Chronic myeloid leukemia (CML) is a malignant myeloproliferative disease characterized by Philadelphia (Ph) chromosome produced by reciprocal translocation between chromosomes 9 and 22 [t (9,22)] (1). The Ph chromosome can be transcribed and encoded into BCR-ABL oncprotein, a constitutive tyrosine kinase that promotes abnormal cellular proliferation and disease progression by stimulating multiple downstream signaling pathways (2). Imatinib (IM), a tyrosine kinase inhibitor (TKI) that target the ATP-binding site of BCR-ABL kinase, has become first-line treatment and effectively improved the survival for the majority of CML patients (3). However, 20-30% of CML patients will develop resistance to IM (4). Hence, there is a desperate need to discover new diagnosis biomarkers of CML and potential targets for CML therapeutics.

Exosomes are nanometer-sized extracellular vesicles (EVs) which are approximately 40-160 nm in diameter (average ~100 nm) (5). When endosomes or multivesicular bodies (MVBs) fuse with the plasma membrane, the exosomes are released (6). The role of exosomes was initially thought to be the carrier of unnecessary cellular waste that remove excess constituents from cells to maintain cellular homeostasis (7). Currently, increasing studies have reviewed that exosomes can carry many functional molecules including proteins, lipids, and nuclear acids (DNA, mRNA, lncRNA, miRNA, and circRNA) derived from the parent cell (8), which serve an essential role in regulating intercellular communication during different physiological and pathological processes (9). In addition, various substances enclosed within exosomes are wrapped by the lipid bilayer, which is relatively stable in body fluids (blood, ascites, saliva, urine, and so on) (10). Due to these unique features of exosomes, they have become ideal candidate biomarkers for the diagnosis, prognosis, and response to therapy of many malignant tumors.

A growing body of research has shown that the exosome is an important part of the tumor microenvironment implicated in chemoresistance development in cancer (11, 12). As for studies related to IM resistance in CML, our previous study found that exosomal miRNA-365 derived from imatinib-resistant (IM-R) cells can be directly transferred to imatinib-sensitive (IM-S) cells and to confer drug-resistance traits (13). Hrdinova and coworkers reported a detailed characterization of the protein cargo of the exosomes derive from K562 and K562IR cells and found IFITM3, CD146 and CD36 are specific exosomal markers associated with the IM resistance (14). However, the single-cell model of CML may not fully reflect the complexity and heterogeneity of CML in vivo. Therefore, we conceive that it is necessary to identify plasma exosomal proteins for finding of reliable biomarkers and potential therapeutic targets associated with the IM resistance in CML.

In this study, we acquired a detailed proteomic profile of exosomes derived from plasma of IM-R and IM-S CML patients. Differentially expressed proteins (DEPs) identified were associated with ribosome, acute myeloid leukemia, prostate cancer, and cholesterol metabolism. We subsequently screened multiple potential biomarkers and verified several in independent cohorts. Our study might be useful for further studies of novel diagnostics and therapeutics for chemoresistance of IM in CML patient.

Conclusion: Proteomic analysis of plasma exosomes provides new ideas and important information for the study of IM resistance in CML. Especially the exosomal proteins (RPL13 and RPL14), which may have great potential as biomarkers of IM resistance.

Keywords: exosomes, chronic myeloid leukemia, proteomics, imatinib, ribosomal protein L13, ribosomal protein L14

Abbreviations: IM, imatinib; TKI, tyrosine kinase inhibitor; CML, chronic myeloid leukemia; LC-MS/MS, liquid chromatograph–mass spectrometer; WB, western blot; PRM, parallel reaction monitoring; DEPs, differentially expressed proteins; Ph, Philadelphia; EVs, extracellular vesicles; MVBs, multivesicular bodies; IM-R, imatinib-resistant; IM-S, imatinib-sensitive; UC, ultracentrifuge; NTA, nanoparticle tracking analysis; TEM, Transmission Electron Microscope; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; GO, Gene Ontology; KEGG, Genes and Genomes.

MATERIALS AND METHODS

Patients and Plasma Samples

All patients were recruited from the Second Affiliated Hospital of Nanchang University (Nanchang, China). The diagnoses, treatment responses, and relapse criteria of CML were made according to the criteria of World Health Organization (15) and the guidelines for diagnosis and treatment of chronic myelogenous leukemia in China (2016 edition) (16). The subjects enrolled in the current study were from newly diagnosed CML patients and none of the patients had received prior TKI therapy or other medical treatment of CML before plasma samples were obtained. Pooled plasma samples from 9 IM-R and 9 IM-S CML patients were used for discovery proteomics analysis. Then, plasma samples from 18 IM-R and 18 IM-S CML patients were used to validate candidate markers. Detailed information of all participants is given in Supplementary Table S1. Peripheral venous blood sample from every single participant was collected in the fasting state and plasma was separated by centrifugation at 2,000 × g for 15 min. The plasma supernatant without lipemia or hemolysis was collected and stored at -80 °C for exosome isolation. This study was conducted in accordance with the Declaration of Helsinki and was approved by the Medical Ethics Committee of the Second Affiliated Hospital of Nanchang University. All patients provided written informed consent.

Exosome Isolation

We isolated exosomes from plasma samples through ultracentrifugation as in our earlier report with slight modifications
The experimental workflow is shown in Figure 2A. The plasma samples were diluted twice with PBS to reduce the viscosity. The mixed liquid was centrifuged to remove cells and cell debris by 10 min centrifugation at 400 × g (4°C), followed by 20 min centrifugation at 2000 × g (4°C). Subsequently, the resultant supernatant was collected and centrifuged at 10,000 × g for 30 min (4°C) and then was filtered through a 0.22 µm syringe membrane filter (Millipore, USA). The supernatant was transferred into ultracentrifuge (UC) tubes with a total volume of 10 ml and ultracentrifuged at 110,000 × g (4°C) for 70 min (Optima-L-80XP, SW41 rotor, Beckman Coulter, USA) to remove debris. Then, the supernatant was gathered and measured with BCA protein quantification kit. To digest proteins, proteins were first reduced by 5 mM dithiothreitol (56°C, 30 min), followed by alkylated with 11 mM iodoacetamide (room temperature, 15 min in darkness). Furthermore, the sample was diluted to a final urea concentration to < 2M by adding 100 mM triethylammonium bicarbonate buffer (TEAB). Finally, proteins were digested twice with trypsin, the first digestion were performed with trypsin (overnight, 37°C) at a ratio of 1:50 (enzyme:protein) and 1:100 for the second digestion (4h).

Exosomal Protein Extraction and Digestion

Exosomes were solubilized in lysis buffer (8 M urea and 1% protease inhibitor) and sonicated three times at high intensity using an ultrasonic processor (Scientz). The suspension was centrifuged at 12,000 × g (4°C, 15 min) to remove debris. Then, the supernatant was gathered and measured with BCA protein quantification kit. To digest proteins, proteins were first reduced by 5 mM dithiothreitol (56°C, 30 min), followed by alkylated with 11 mM iodoacetamide (room temperature, 15 min in darkness). Furthermore, the sample was diluted to a final urea concentration to < 2M by adding 100 mM triethylammonium bicarbonate buffer (TEAB). Finally, proteins were digested twice with trypsin, the first digestion were performed with trypsin (overnight, 37°C) at a ratio of 1:50 (enzyme:protein) and 1:100 for the second digestion (4h).

Liquid Chromatography-MS/MS Analysis

Digested samples were subjected to LC-MS/MS detection using a minor modifications of previously described method by Hou et al. (18). The tryptic peptide mixture was resuspended in solvent A (0.1% formic acid) and directly inject into a home-made reversed-phase analytical column (15 cm length, 75 µm i.d.). Then, the peptides were separated on an EASY-nLC 1000 UPLC system (Thermo Fisher Scientific) with a linear gradient mobile phase buffer at constant flow rate of 400 nL min⁻¹. The chromatography gradient was carried out starting at 6% solvent B (0.1% formic acid in 98% acetonitrile), increased to 23% B over 26 min, changed to 35% in 8 min, climbing to 80% in 3 min and maintenance at 80% for an additional 3 min. The eluted peptides were loaded into the nanospray ionization (NSI) source and performed tandem mass spectrometry (MS/MS) analysis on a Q Exactive™ Plus (Thermo Fisher Scientific) coupled online to the UPL (Thermo Fisher Scientific). The applied voltage of electrospray was 2.0 kV. The intact peptides were detected for MS full scans with the scanning range of m/z 350 to1800 and a resolution of 70,000 in the Orbitrap. Then, the selected peptides were detected at a resolution of 17,500 and normalized collision energy (NCE) of 28. The MS analysis was performed by a data-dependent scan mode that alternated between one MS scan and 20 MS/MS scans, and the dynamic exclusion was set to 15.0 s. Automatic gain control (AGC) target value and fixed first mass were set to 5E4 and 100 m/z, respectively.

Database Search

The secondary mass spectrum data were retrieved using Proteome Discoverer (PD) 2.4 software (Thermo Scientific) with the Maxquant search engine (v1.5.2.8). In detail, the raw MS/MS data files were searched against the human UniProt/SwissProt proteome database with reverse decoy database. The cleavage enzyme was set as Trypsin/P, and the number of missed cleavages was allowed up to two. The length of minimal peptide specified to be 7 amino acid residues, the maximal modification number of allowed per peptides was set as 5. Precursor mass
tolerance were set to 10 ppm and 5 ppm for the first search and the main search, respectively. The fragment ion mass tolerance was as 0.02 Da. The acetylation modification on N-terminal and oxidation modification on methionine were designated as variable modifications. The carbamidomethyl modification on cysteine residues was designated as a fixed modification. The quantitative method of proteins was label-free quantification (LFQ), and the false discovery rate (FDR) of protein identification and peptide-spectrum match (PSM) identification was set as 1%.

Bioinformatics Analyses
Gene Ontology (GO) annotation proteome was obtained based on the UniProt-GOA database (http://www.ebi.ac.uk/GOA/). The fold change values ≥ 1.5 for the upregulated proteins and ≤ 0.667 for the downregulated, which were used as the criteria to identify the DEPs between IM-R and IM-S samples. We performed the bioinformatics analyses in order to further analyze the biological function of DEPs. For GO functional enrichment analysis, proteins were categorized into three ontologies (biological process, molecular function and cellular compartment) using the extension R package ‘ClusterProfiler’ (19)(https://www.bioconductor.org/packages/release/bioc/html/ClusterProfiler.html). We used the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (http://www.genome.jp/kegg/) to perform pathway enrichment analysis. The enrichment of DEPs was detected by double-tailed Fisher’s exact test. The adjusted p-value <0.05 for enrichment analysis was considered significant. Protein interaction networks were analyzed by STRING (20) (https://string-db.org/v.11.0), and the parameter settings were as follows: the minimum required interaction scores were set at highest confidence (0.900), the network display options were set at hide disconnected nodes in the network, the other parameters were set to default values. The essential hub proteins screens were performed by Cytoscape software (21).

PRM-MS Analysis
Twenty proteins were screened through bioinformatics analyses, and further verified with PRM. The digested peptides were dissolved in 0.1% formic acid (mobile phase A) and analyzed using PRM method on Exploris 480™ mass spectrometer (Thermo) coupled to an Easy-NLC 1200 UPLC-system (ThermoFischer). The detection and analysis of peptide parent ions and their secondary fragments were performed in the Orbitrap analyzers. Data was acquired with Targeted Mass modes. Higher-energy collision dissociation (HCD) was set at normalized collision energy of 27%. Mass spectrometer settings were, for full MS acquisitions, mass-to-charge ratio (m/z) range set at 395–815 m/z, resolution set at 60,000, automatic gain control (AGC) set at 300%, maximum ion injection time of 100 ms; for MS/MS acquisitions, resolution set at 15,000, automatic gain control (AGC) set at 75%, maximum ion injection time set to 100 ms, isolation window set to 1.6 m/z.

Statistical Analyses
All other data were analyzed using GraphPad Prism (version 8.0.2; La Jolla, CA, USA). The Mann-Whitney U-test was used to compare differences of the exosomal RPL13 and RPL14 levels between IM-R and IM-S CML patients. The values of p < 0.05 (two-sided) were considered statistically significant.

RESULTS

General Experiment Design
We designed a simple workflow to understand the protein composition of the CML plasma exosomes and to screen potential diagnostic exosomal protein biomarkers for IM-R CML patients, as illustrated in Figure 1. For proteomics analysis, plasma samples obtained from 9 IM-R and 9 IM-S CML patients were pooled, respectively. Exosomes were isolated from the pooled plasma samples using ultracentrifugation method and verified using TEM, NTA, and WB. The gained exosomes were lysed to extract proteins and were digested into peptides by trypsin. The exosomal protein profiles were analyzed by label free quantitative proteomics based on LC-MS/MS. DEPs and candidate protein biomarkers were determined by bioinformatics analyses, and then PRM analyses were applied to validate the candidate proteins. Based on the validation results, two proteins were selected as diagnostic markers and the expression levels of the selected proteins were further validated by WB in 18 IM-R and 18 IM-S CML patients.

Characterization of Plasma Exosomes
Figures 2B–D show representative images of plasma exosomes obtained with TEM, NTA, and WB. NTA results revealed the average size of the exosomes was approximately 100 nm in diameter (Figure 2B). TEM demonstrated that the exosomes took on cup-shaped membrane vesicles and the sizes consistent with the NTA results (Figure 2C). We further confirmed the presence of characteristic exosomal membrane markers including ALIX, TSG101, and CD81 by WB. Altogether, these results illustrated that exosomes from clinical plasma samples were successfully collected and purified.

Proteomic Profiling of Exosomes From IM-R and IM-S CML Patients
In order to achieve the characteristics of exosome proteome in the IM-R and IM-S CML patients, protein profiling was performed on the extracted plasma exosomes. 14990 unique peptides were identified from plasma exosomes in all groups, representing 2812 proteins, including 2729 quantifiable proteins (Figure 3A and Supplementary Table 2). Principal component analysis of the data revealed a group of data (CML-S1) in the sensitive group with a large degree of dispersion (Figure 3B), and this group was removed during further analysis. As shown in Figure 3C and Supplementary Table 3, a total of 2654 quantifiable proteins were identified in all groups. Among them, 2276 proteins (85.8%) overlapped between the IM-R and IM-S group, 353 (13.3%) and 25 (0.9%) proteins were found in IM-R and IM-S group, respectively. In order to verify the relationship between the plasma exosomal proteins and known vesicular proteins, we compared quantified exosomal proteins from the two plasma groups with the proteins in Exocarta (22,
23) and Vesiclepedia databases (24, 25). The result was displayed as Venn diagram in Figure 3D. Of the 2654 identified plasma exosomal proteins, 2005 (75.55%) proteins were shared by the three data sets, and 234 (8.82%) proteins are only present in our study, which may be newly discovered exosomal proteins. In addition, out of the top 100 reported extracellular vesicle markers in ExoCarta, and Vesiclepedia database, 73 proteins were identified in our study (Figure 3E). These data confirmed that the exosomes we prepared are rich in exosomal proteins.

**GO Classification and KEGG Pathway Analysis of DEPS**

In order to explore the difference of exosomal proteins between the IM-R group and the IM-S group, a fold change ≥1.5 or ≤0.667 and p value <0.05 was used as a threshold to identify DEPs. As illustrated in the volcano plot and heat map (Figures 4A, B, and Supplementary Table 4), we screened 279 DEPs under this condition, including 151 upregulated proteins and 128 downregulated proteins. To analyze the biological function of the DEPs further, we conducted GO Term and KEGG pathway enrichment analysis. GO analysis results were shown in Figures 4C, D. Cellular component (CC) analysis revealed that the upregulated proteins were enriched in cytosolic part, cytosolic ribosome, ribosome, and ribosomal subunit, whereas the downregulated proteins were enriched in extracellular region part, extracellular space, and lipoprotein particle. As for molecular functions (MF), the upregulated proteins were associated with structural constituent of ribosome, whereas the downregulated proteins were associated with lipid transporter activity. As for biological processes (BP), the upregulated proteins were involved in translation, amide biosynthetic process, and peptide biosynthetic process, whereas the downregulated proteins were involved in protein-lipid complex remodeling, protein-lipid complex assembly, and cholesterol transport. Through KEGG pathway annotations, we found that ribosome, acute myeloid leukemia and prostate cancer were the most enriched pathways involved in the upregulated proteins (Figure 4E), while the downregulated proteins were enriched in pathways associated with cholesterol metabolism (Figure 4F). The functional analysis showed that the upregulated and downregulated proteins were related to ribosome and lipid metabolism, respectively. More detailed results of GO Term
and KEGG pathway enrichment analysis can be obtained from Supplementary Table 5.

Interaction Network Construction of DEPs and Identification of Key Proteins

To investigate the interactions of DEPs, we constructed a comprehensive network using the STRING database with high confidence expressions (scores >0.9) and visualized with Cytoscape 3.8.1. In total, there are 155 DEPs in the PPI network, which contains 83 upregulated and 72 downregulated proteins (Figure 5A). Next four different algorithms (MCC, MNC, DMNC, and Degree) of the cytoHubba plugin were used to screen the hub proteins (Figures 5B–E) and 16 proteins are the common hub proteins (Figure 5F). Figure 5G is a venn diagram consisting of three groups, 62 proteins in the red box are present in every pooled IM-resistant CML samples and are not present in IM-sensitive CML samples. Finally, 16 common hub proteins overlapped with 62 proteins expressed only in the RR group and 4 proteins (RPL14, RPL18A, RPS15A, and RPL13) were screened to as candidate proteins, as shown in Figure 5H.

Verification of Candidate Biomarkers with PRM and Western Blot Analysis

To validate the candidate proteins and the MS results, we selected twenty proteins including four candidate proteins for quantification using PRM. Of the 20 proteins, 13 proteins were detected and the fold difference was consistent with label-free quantification. In particular, the candidate proteins RPL14, RPS15A, and RPL13 were only identified in IM-R CML patients, and not in IM-S CML patients. Based on the fold change as well as the result of PRM, candidate exosomal proteins
RPL14, and RPL13 were selected for validation by WB analysis in individual samples (18 IM-R and 18 IM-S CML patients). The exosomal standard marker TSG101 was used as the internal control standard. The levels of RPL14, and RPL13 were significantly higher in IM-R CML patients compared with IM-S CML patients, which was consistent with proteomics and PRM results. The results are shown in Figure 6 and Table 1.

DISCUSSION

The development of IM resistance is a significant clinical challenge for the treatment of CML (26). Fully understanding the mechanisms of resistance to IM will help us to overcome this problem in the future. The point mutation or amplification of \textit{BCR-ABL} gene are the main resistant mechanisms to IM (27). Meanwhile, growing studies have shown that exosomes were linked with drug resistance (28, 29), by which exosomes from chemoresistant cells can transfer resistance to chemosensitive cells in a variety of tumors (30, 31), including CML (13, 14).

Furthermore, Exosomes contain various specific substances from target cells and are stable in various bodily fluids including blood, urine, saliva, ascites fluids, and cerebrospinal, making them ideal biomarkers for liquid biopsies. Multiple studies have shown that exosomal protein markers with diagnostic significance of chemoresistance have been obtained in different tumors such as Prostate Cancer (32, 33), Breast Cancer (34), and B Cell Lymphoma (35). One study demonstrated that exosomes released by MSCs transferred their proteasome inhibitors resistance phenotype to the multiple myeloma cells, through PSMA3 and PSMA3-AS1 present in the cargo of exosome. Plasma exosomal proteins PSMA3 and PSMA3-AS1 might be a promising therapeutic targets and prognostic predictors for proteasome inhibitors resistance (36). Moreover, hypoxia-induced exosomes contain PKM2 protein, which were able to modulate resistance to cisplatin. Exosomal PKM2 may serve as a promising predictive biomarker for cisplatin treatment response in NSCLC (37). However, the plasma exosomes proteome in drug resistance of CML have rarely been investigated. Here, we performed a comparative proteomic analysis of the plasma
exosomes from IM-R and IM-S CML patient in order to identify proteins that may have a relevant role for IM resistance in CML.

In this study, we first isolated plasma exosomes and comprehensively characterized them using multiple methods. The TEM results shown that the morphological structure and size of plasma exosomes were consistent with the typical exosomes. Western blot revealed that the representative markers of exosomes, such as CD81, TSG101, and ALIX were detected in plasma exosomes. These results shown that we successfully isolated exosomes from plasma of IM-R and IM-S CML patients by ultracentrifugation. Furthermore, we established a solid foundation to facilitate further investigations of the exosome proteomic of CML. Then we carried out proteomic study on the plasma exosome by label free quantitation based on LC-MS/MS and identified IM-resistance associated potential biomarkers. The proteome analysis revealed 2654 quantifiable proteins in all groups. Out of the total 2654 proteins, 2005 (75.55%) proteins were found in the ExoCarta and Vesiclepedia databases, providing further evidence for the identity of isolated plasma exosomes. In addition, 234 (8.82%) proteins are not present in the ExoCarta and Vesiclepedia databases, which may be newly discovered exosomal proteins. Certainly, this assumption needs to be further validated. There were 279 DEPs with 151 upregulated and 128 downregulated proteins in IM-R CML patients compared with IM-S CML patients. The GO and KEGG functional analysis shown that the upregulated and downregulated proteins were related to ribosome and lipid metabolism, respectively. These DEPs were helpful in illustrating the resistance mechanisms of IM in CML. To screen resistance biomarker to IM in CML, we further analyzed the DEPs. Sixteen hub proteins were screened from DEPs through the PPI network and the cytoHubba plugin of Cytoscape software. Among the 16 hub proteins, 4 proteins (RPL14, RPL18A, RPS15A, and RPL13) were expressed only in
the drug-resistant group and selected as candidate biomarker for further studies. PRM verification analysis is a high throughput quantitative technique with high accuracy (38). Twenty proteins including RPL14, RPL18A, RPS15A, and RPL13 were selected for further verification by PRM quantification. 13 out of 20 proteins were detected and the expression trend was consistent with the results of MS, indicating the good reliability of the proteomic profiling. Finally, we selected two proteins for validation by WB in an independent cohort with 18 resistant and 18 sensitive CML. Surprisingly, the results confirmed that exosomal RPL13 and RPL14 levels in IM-R CML patients were significantly higher than IM-S CML patients.

Ribosomal proteins are central components of the ribosome and are involved in protein biosynthesis. However, extraribosomal functions of ribosomal proteins were gradually recognized (39, 40). Recent studies have found that ribosomal proteins were closely associated with tumorigenesis (41), tumor development (42), and resistance to antitumor drugs (43). For example, the expression of RPL34 was found to be more highly in osteosarcoma tissues compared with normal adjacent tissue and was correlated with a poor prognosis (44). RPL41 can induce the degradation of the activating transcription factor 4 (ATF4) to regulate tumour cell death, cell cycle, and chemosensitivity. Therefore, RPL41 may have potential as an anti-ATF4 agent for cancer therapy (45).
These studies strongly support that the ribosomal protein might act as biomarkers and therapeutic targets for tumor.

RPL13 and RPL14 belong to ribosomal proteins and are a component of the large subunit of the ribosome. RPL13 was highly expressed in gastrointestinal cancer and enhanced the proliferation of gastrointestinal cancer cells and may be a useful molecular marker of drug efficacy in gastrointestinal cancer (46). In kidney renal clear cell carcinoma (KRCCC), researchers found that RPL13 was associated with the prognosis of tumor patients (47). Huang found that RPL14 are frequently altered during tumorigenesis of esophageal squamous cell carcinomas (ESCC) and could be used for the early detection of ESCC (48). In addition, RPL14 may be a tumor suppressor gene in nasopharyngeal carcinoma (NPC), which could inhibit cancer progression (49). Bioinformatics analyses suggests that the low expressions of ribosomal protein L14 (RPL14) indicated poor
prognosis in Triple-negative breast cancer (TNBC) (50). In contrast to the above studies, some studies have shown that RPL14 may be an oncogenes, down-regulation of RPL14 could inhibit the development of cervical cancer (51), and high expression of RPL14 could promote the migration and invasion of cervical cancer cells (52). Thus, ribosomal proteins may play different functions in different tumors. In our study, we found that plasma exosome proteins RPL13 and RPL14 were highly expressed in IM-R CML patients, and could be used as potential markers for predicting and monitoring IM response in CML patients. However, whether it was involved in IM resistance in patients with CML and its corresponding mechanism remains to be further confirmed.

There are some unavoidable flaws in our study. For example, the exosomes in plasma are derive from various cells in the human body, including tumor cells and non-tumor cells. Therefore, the exosomes we extracted from patient plasma were non-tumor-specific, with strong heterogeneity from various cell. Simultaneously, due to the limitations of extraction methods of exosomes, we cannot exclude the possibility of the extracted exosomes being mixed with contaminated proteins or other particles. These heterogeneous exosomes or contaminants may have some impact on the experimental results, which are unavoidable problems (common problems) in the current field of exosome research. On the other hand, the cohort used to verify marker proteins was a relatively small number of patients from a single center, which is not enough to ensure the diagnostic efficacy of the screened marker proteins. Our results will need further verification with large samples from multi-centers in the future.

CONCLUSION
In summary, we successfully isolated and characterized plasma exosomes from CML patients. Meanwhile, we also first identified and compared the global proteomic profiles of plasma exosomes between IM-R and IM-S patients. Four significant proteins were screened and two of them (RPL13 and RPL14) were validated as novel potential biomarkers of IM-resistance in CML. These findings will provide new ideas and important information for further studies of IM resistance in CML.

DATA AVAILABILITY STATEMENT
The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: PRIDE under accession number is PXD028779.

ETHICS STATEMENT
The studies involving human participants were reviewed and approved by the Medical Ethics Committee of the Second Affiliated Hospital of Nanchang University. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS
BH and X-ZW designed the study. M-YL, CZ, LC, F-YY, Q-HM, JLin, H-BZ, and JLIu were responsible for the collection of the clinical samples. M-YL, CZ, YC, SX, and Y-LY performed the experiments. M-YL, F-MZ, and J-YJ analyzed the data. M-YL wrote the main manuscript. BH and X-ZW contributed to manuscript review and revision. BH and X-ZW contributed equally to this work. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2021.779567/full#supplementary-material

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