Comparative Study of Three Mixing Methods in Fusion Technique for Determining Major and Minor Elements Using Wavelength Dispersive X-ray Fluorescence Spectroscopy

Dan-Ping Zhang 1,2,3, Ding-Shuai Xue 3,*, Yan-Hong Liu 3, Bo Wan 3, Qian Guo 3 and Ju-Jie Guo 3

1 State Key Laboratory of Lithospheric Evolution, Institute of Geology and Geophysics, Chinese Academy of Sciences, Beijing 100029, China; zdanp@mail.iggecas.ac.cn
2 Innovation Academy for Earth Science, Chinese Academy of Sciences, Beijing 100029, China
3 Institute of Geology and Geophysics, Chinese Academy of Sciences, Beijing 100029, China; liuyanhong@mail.iggecas.ac.cn (Y.-H.L.); wanbo@mail.iggecas.ac.cn (B.W.); guoqian@mail.iggecas.ac.cn (Q.G.); guojieju@mail.iggecas.ac.cn (J.-J.G.)

* Correspondence: xuedingshuai@mail.iggecas.ac.cn; Tel.: +86-010-8299-8487

Received: 13 August 2020; Accepted: 7 September 2020; Published: 17 September 2020

Abstract: Accurate analysis using a simple and rapid procedure is always the most important pursuit of analytical chemists. In this study, a new sample preparation procedure, namely the shaker cup (SH) method, was designed and compared with two sample preparation procedures, commonly used in the laboratory, from three aspects: homogeneity of the sample–flux mixture, potential for sample contamination, and sample preparation time. For the three methods, a set of 54 certified reference materials (CRMs) was used to establish the calibration curves, while another set of 19 CRMs was measured to validate the results. In the calibration procedures, the matrix effects were corrected using the theoretical alpha coefficient method combined with the experimental coefficient method. The data of the major oxides (SiO₂, TiO₂, Al₂O₃, Fe₂O₃, MnO, MgO, CaO, Na₂O, K₂O, and P₂O₅) and minor elements (Cr, Cu, Ba, Ni, Sr, V, Zr, and Zn) obtained by wavelength dispersive X-ray fluorescence spectroscopy (WD-XRF) were compared using two derivative equations based on the findings by Laurence Whitty-Léveillé. The results revealed that the WD-XRF measured values using the SH method best agreed with the values recommended in the literature.

Keywords: wavelength dispersive X-ray fluorescence spectroscopy; sample preparation procedure; fused glass disc; shaker cup method

1. Introduction

Wavelength dispersive X-ray fluorescence spectrometry (WD-XRF) is a widely used qualitative and quantitative analytical method, which has been applied in many fields including geology, geochemistry, mining, metallurgy, non-ferrous metals, building materials, environmental protection, and commodity inspection [1–8]. In geology and geochemistry, WD-XRF is mainly used to analyze the elemental composition of rocks, which can be employed for the exploration of mineral deposits and classification of the whole rock diversity [9,10]. Atomic absorption spectrometry (AAS), inductively coupled plasma atomic emission spectrometry (ICP-AES), and inductively coupled plasma mass spectrometry (ICP-MS) are also used for the analysis of major, minor, and trace elements in rocks; however, these instrumental methods require tedious wet chemical pretreatment. Thus, burdensome labor can easily cause accidental contamination and uncertainty of analysis, while many reagents that are toxic to the
operator are consumed [11]. In contrast, XRF presents numerous advantages including simple sample preparation, no environmental pollution, and little training for personnel. This technique can also detect over a wide range of elements and can be employed to analyze multiple elements simultaneously. Moreover, XRF instruments have long-term stability enabling sample analysis with high precision and reproducibility. To take full advantage of these features, however, one needs to pay particular attention to the sample preparation process. This is because such processes have a very high potential for introducing systematic errors into the XRF analysis.

The most common sample preparation methods used for the analysis of rock samples by conventional WD-XRF instrumentation are pressed powder pellets and fused glass discs [12,13]. Fused glass discs, which is the most extensively used analytical technique in the laboratory, is excellent for analyzing major and minor elements as it reduces matrix effects, eliminates particle size effects, and provides a homogeneous specimen [14,15]. Many studies to determine the major and minor elements using WD-XRF have been reported in the literature [9,10,16–18]. The results have revealed that 1:10 dilution glass discs are best suited for major and minor element analysis as this ratio accounts for the sample, reagent dosage, analysis signal strength, and simplicity and feasibility of the operation procedure. Preferably, the sample should be homogeneously mixed with flux before it is fused to a glass disc. Thus, different sample preparation procedures for making fused glass discs have been reported. For example, Shintaro Ichikawa and Toshihiro Nakamura mixed the sample and flux on paraffin paper using a bamboo spatula and subsequently placed the mixture into a Pt–Au crucible for fusing [19]. Masatsugu Ogasawara et al. also used weighing paper and mixed the sample and flux thoroughly with a spatula. Additionally, they suggested another alternative in which the sample and flux powder are transferred to an agate mortar, mixed thoroughly, and subsequently transferred into a Pt–Au crucible for fusing [20]. Atsushi Goto and Yoshiyuki Tatsumi reported a mixing procedure in which the powder rock sample was transferred from the weighing paper onto the flux and the two were then mixed by raising the sides of the weighing paper; however this technique is difficult to operate and the sample can be easily spattered [16]. Kenichiro Tani et al. used a vibration touch-mixer to mix the sample and flux, which were both directly weighed into the same Pt–Au crucible [21]. Yuji Orihashi and Takafumi Hirata also reported mixing using a touch-mixer [22]. However, heterogeneous glass discs made by this method have only been successfully used to mix an aqueous solution in which convection occurs effectively, which is not observed in solid powders. W. B. Stern reported a method in which an ignited powder sample was carefully mixed in an agate mortar with dried Li$_2$B$_4$O$_7$ and subsequently transferred into a Pt–Au crucible for fusing [23]. However, these mixing methods were not described in enough detail. Moreover, many authors such as Toru Yamasaki [24] and Atsushi Kamei [25] did not even report the mixing method employed.

In view of the XRF analysis error mainly originating from the sample preparation procedure, this step is extremely important for the whole XRF analytical process [26]. Therefore, it is essential to explore a simple and effective way to prepare a homogeneous glass disc, prior to XRF analysis, from which more accurate and precise analytical results can be obtained.

In our lab, two sample preparation procedures, namely the grinding (GR) and stirring rod (ST) methods, have been mainly used to prepare fused glass discs. The GR method comprises mixing of the sample and flux in an agate mortar and subsequent transfer of the afforded mixture to a Pt–Au crucible with a spoon and brush for the next fusing step; this method was also used by Stern [23]. The advantage of this method is that the sample and flux can be mixed very evenly, whereas its main disadvantage is the ease of contamination and sample loss during the transfer process. The ST method comprises the direct weighing of the sample and flux into a Pt–Au crucible, mixing with a glass rod, and subsequent fusion of the resultant mixture in a fusion machine. The advantage of this method is that it is a simpler process with a lower risk of contamination and sample loss during the transfer process. On the other hand, its main disadvantage is insufficient mixing of the sample and flux.

Herein, we propose an improved sample preparation procedure, namely the shaker cup (SH) method, inspired by the protein shaker cup used in bodybuilding. This method combines the
advantages of the GR and ST methods to conveniently mix the sample and flux homogenously in a cheap shaker cup. Moreover, this method allows easy transfer of the afforded mixture to a Pt–Au crucible. To evaluate the quality of the data obtained from the different glass disc procedures, calibration curves were established and major and minor elements of 19 certified reference materials (CRMs) were examined. The operational processes, lower limits of detection (LLD), accuracy, and precision were compared in detail. The results revealed that of the three investigated methods, the SH method is the most suitable for creating fused glass discs.

2. Experimental

2.1. Instrumentation

An electric drying oven (WGL-125B, Tianjin Taisite Instrument Co., Ltd., Tianjin, China) was used to dry the samples and flux, while a muffle furnace (SX-GO4133, Tianjin Zhonghuan Test Electrical Furnace Co., Ltd., Tianjin, China) was used to burn the samples. The samples and flux were weighed with an electronic balance (ME104E, Mettler-Toledo, Zurich, Switzerland). An ultrasonic cleaning device (Elma E100H, Elmasonic, Singen, Germany) was used to clean the Pt–Au crucibles and molds, while a six-position automatic fusion device (Claisse Fluxy, Corporation Scientific Claisse Inc., Quebec, Canada) was used to create the fused glass discs. Imaging studies were performed on an energy dispersive X-ray fluorescence spectroscopy (EDXRF) spectrometer (M4 TORNADO, Bruker Nano GmbH, Berlin, Germany). The X-ray fluorescence spectrometer (AXIOS, PANalytical, Almelo, The Netherlands) was used with a Rh anode X-ray tube and 4 kW excitation power. This is a sequential instrument with a single goniometer-based measuring channel covering the complete elemental measurement range from F to U, in the concentration range from 1.0 ppm to % level determined in vacuum media. SuperQ 4.0 analysis software was used. The spectrometer conditions for ten major and eight minor elements are listed in Table 1.
Table 1. Instrumental parameters used in wavelength dispersive X-ray fluorescence spectrometry (WD-XRF).

| Element | Si | Ti | Al | Fe | Mn | Mg | Ca | Na | K | P | Cr | Cu | Ba | Ni | Sr | V | Zr | Zn |
|---------|----|----|----|----|----|----|----|----|---|---|----|----|----|----|----|---|---|---|
| Line    |    |    |    |    |    |    |    |    |   |   |    |    |    |    |    |   |   |   |
| Crystal | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka | Ka |
| Detector| Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow | Flow |
| Voltage/kV | 30 | 40 | 30 | 60 | 30 | 60 | 60 | 60 | 40 | 60 | 40 | 60 | 60 | 60 | 60 | 60 | 60 | 60 |
| Intensity/µA | 120 | 90 | 120 | 60 | 60 | 60 | 60 | 60 | 60 | 90 | 60 | 60 | 60 | 60 | 60 | 60 | 60 | 60 |
| Collimator/µm | 150 | 300 | 600 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 | 300 |
| Counting time/s | 60 | 30 | 40 | 26 | 36 | 40 | 40 | 40 | 30 | 30 | 40 | 40 | 40 | 40 | 40 | 40 | 40 | 40 |
| Peak Angle/° | 109.1896 | 86.1198 | 144.7838 | 57.5056 | 62.9604 | 23.2312 | 113.0678 | 28.0754 | 136.6656 | 141.1432 | 69.3396 | 45.0206 | 87.1566 | 48.6566 | 25.1452 | 76.5228 | 22.5018 | 41.7962 |
| Bg1 | 3.3508 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 |
| Bg2 | 3.3508 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 | 3.1288 |
| PHD | 32–69 | 37–63 | 33–55 | 30–72 | 35–65 | 30–70 | 35–65 | 30–70 | 35–65 | 30–70 | 35–65 | 30–70 | 35–65 | 30–70 | 35–65 | 30–70 | 35–65 | 30–70 |
| Peak Angle/° | 109.1896 | 86.1198 | 144.7788 | 57.5038 | 62.9598 | 23.2312 | 113.0678 | 28.0600 | 136.6650 | 141.1414 | 69.3388 | 45.0214 | 87.1556 | 48.6578 | 25.1452 | 76.9202 | 22.5018 | 41.7962 |
| Bg1 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 | 2.2642 |
| Bg2 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 | 3.1132 |
| PHD | 31–72 | 34–66 | 33–53 | 29–75 | 32–70 | 23–66 | 31–69 | 26–58 | 37–62 | 35–52 | 33–53 | 34–53 | 32–78 | 32–69 | 24–78 | 32–53 | 32–78 | 25–70 |
| Peak Angle/° | 109.1896 | 86.1226 | 144.7788 | 57.5018 | 62.9538 | 23.2312 | 113.0678 | 28.0708 | 136.6650 | 141.1414 | 69.3388 | 45.0214 | 87.1576 | 48.6570 | 25.1452 | 76.5258 | 22.5018 | 41.7962 |
| Bg1 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 | 2.3086 |
| Bg2 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 | 3.1338 |
| PHD | 30–71 | 35–65 | 22–78 | 36–64 | 34–52 | 29–75 | 32–69 | 24–58 | 31–71 | 26–60 | 37–63 | 36–52 | 33–53 | 34–52 | 33–53 | 34–52 | 33–53 | 34–52 |

The Grinding Method

The Stirring Rod Method

The Shaker Cup Method
2.2. Reagents

Absolute ethanol (Sinopharm Chemical Reagent Co., Ltd., Shanghai, China) was used for scrubbing the crucibles and agate mortar. Citric acid monohydrate (Sinopharm Chemical Reagent Co., Ltd., Shanghai, China) was used to clean the Pt–Au crucibles and molds, while ammonium bromide (Sinopharm Chemical Reagent Co., Ltd., Shanghai, China) was used as an exfoliation agent. The ultra-pure grade mixed reagent lithium tetraborate: lithium metaborate (67:33; Claisse, Quebec, Canada) was used as an alkali flux to create the fused glass discs.

2.3. Reference Materials

Fifty-four CRMs were used to calibrate the spectrometer. We considered the quality of the recommended values for each CRM, required interval of concentration for each element, and previous calibration tests. The source and recommended values for these reference materials were obtained from a series of literature. In the calibration procedure, the matrix effects were corrected using a theoretical alpha coefficient method combined with an empirical coefficient method. Nineteen CRMs were used for the validation of the three sample preparation procedures. The details of the 73 CRMs employed in this study are listed in Table S1.

2.4. Procedure

The procedure used to create the fused glass discs from the rock powder samples is illustrated in Figure S1 and the details can be described as follows: first, place the samples and flux into an oven, then dry them at 105 °C for 12 h. Second, take the samples and flux out, then put them into a desiccator to cool for 2 h at 25 °C. Third, weigh 0.6000 g of powered rock sample and 6.000 g of flux. Fourth, mix the weighed sample and flux, then transfer them into a Pt–Au crucible (25 mL; 95% Pt, 5% Au). Fifth, add four drops of 0.12 g mL\(^{-1}\) ammonium bromide solution (0.18 mL) into the Pt–Au crucible, then create six fused glass discs simultaneously with the six-position automatic fusion device. The heating time is 19 min at 1050 °C. The cooling time is 285 s. Last, take out the fused glass discs with a suction pen after fusing, then number each one according to the sample name on the non-test surface and put them into a desiccator for subsequent testing.

Notably, most laboratories tend to employ the same drying, cooling, weighing, and numbering steps, while the mixing step tends to be different. Moreover, the adjustable factors involved in automatic melting using the Claisse Ox fusion furnace are beyond the scope of this article and are not elaborated herein. The procedure flow charts are presented in Figure S2 and the sample preparation methods are described in detail. For the grinding method, first, wipe 30 5 mL porcelain crucibles and 30 25 mL porcelain crucibles with alcohol. Second, place the 5 mL porcelain crucibles into a muffle furnace and burn at 1000 °C for 30 min, then take them out and put them into a dryer to cool for 1 h at 25 °C, then weigh each one. Third, weigh 0.6000 g of different dried sample into each 5 mL porcelain crucible, and 6.0000 g of flux into each 25 mL porcelain crucible. Fourth, put the 5 mL porcelain crucibles with weighed samples into a muffle furnace and burn at 1000 °C for 1 h, then take them out and put them into a desiccator to cool for 2 h at 25 °C, then weigh each one. Last, pour each burned sample into an agate mortar to grind until it no longer has the particle sense, then pour in the weighed flux, fully grind and mix, then transfer the mixture to a Pt–Au crucible using spoon and brush. For the stirring rod method, weigh 6.000 g of flux into a cleaned Pt–Au crucible directly, then weigh 0.6000 g of dried sample into a small aluminum dustpan, then pour it into the Pt–Au crucible filled with weighed flux, and then mix them with glass rod manually. For the shaker cup method, first, weigh 6.000 g of flux into a 40 mL commercial shaker cup which is disposable product, and can be used directly. Second, weigh 0.6000 g of dried sample into a small aluminum dustpan, then pour it into the shaker cup filled with flux, then cover the lid. Last, shake it several times manually, and then pour the mixture into a Pt–Au crucible. Next, add two drops of deionized water into the shaker cup, cover the lid, and shake it...
for several times manually again, then pour the mixture into the Pt–Au crucible to remove powder stuck to the wall.

3. Results and Discussion

3.1. Operational Process

In quantitative XRF analysis, the whole process can be divided into three steps, which constitute the source of the overall analysis error, namely sampling, sample preparation, and spectrometric analysis [27]. Rudolf Muller stated that the errors due to sample preparation are superimposed onto other errors of measurement, thereby increasing the total error [28]. Furthermore, John et al. reported that the overall error is equal to the square root of the sum of the squares of error of the individual components of the error. Thus, because of the good stability of the XRF instrument and experimental and theoretical correction algorithms available in the software, the largest error stems from sample preparation [29]. This indicates that the sample preparation procedure is the most important step in the analysis, which strongly influences the final quantitative result. To explore the possibility of a new superior sample preparation method, three sample preparation methods were compared in the following aspects: homogeneity of the sample–flux mixture, potential contamination and loss during sample preparation, and sample preparation time.

The homogeneity of the fused glass discs from the three different sample preparation methods was determined by observing the color and mixedness (Figure 1) of the sample–flux mixtures and comparing the elemental mapping images. In the GR method (Figure 1a), the sample and flux were mixed very evenly (consistent mixture color) prior to fusing and had no particle sense. In the ST method (Figure 1b), the sample and flux were mixed unevenly and were in a two-phase separation state. Indeed, the particle point (red arrows) indicated the presence of agglomeration in some of these mixtures. Finally, for the sample–flux mixture prepared by the SH method (Figure 1c), the mixing effect was better than that of the mixture prepared by the ST method, and there was no significant two-phase separation state due to the marked impact between the sample and flux in the shaker cup.

![Figure 1. Sample–flux mixtures attained by the (a) grinding (GR), (b) stirring rod (ST), and (c) shaker cup (SH) methods.](image)

The mixture could be further mixed at the fusing step; thus, we next compared the uniformity of the three different sets of fused glass discs. Figure 2 illustrates the mapping images of some elements of GBW07101. The images indicated that there was no significant difference in the homogeneity of the
three methods, and the major and minor elements were very evenly distributed. However, there was a significant difference in the content of volatile elements such as Cl. Indeed, the Cl content in the GR-prepared mixture was significantly lower than that of the mixtures prepared by the other two methods. This was attributed to the muffle furnace burning of the CRMs in the GR method. Notably, although no differences in the uniformity of the fused glass discs were observed in the mapping images, in the actual ST sample preparation process, the fused glass discs often displayed crystallization spots that led to cracking of the fused glass discs (Figure 3). This was mainly attributed to insufficient sample mixing prior to melting and the absence of further decomposition in the fusion machine, which ultimately led to the failure of glass disc formation. Thus, for this method, the mixtures had to be fused two- or three-fold to attain a homogeneous glass disc.

Figure 1. Sample–flux mixtures attained by the (a) grinding (GR), (b) stirring rod (ST), and (c) shaker cup (SH) methods.

Figure 2. Mapping images of the GBW07101 (ultramafic rock) samples [Scanning area: width, 4645 pixels (92.9 mm) and height, 1615 pixels (32.3 mm); counting time per pixel (20 μm), 5 ms].

The potential sample contamination and losses during sample preparation were significant for all three preparation methods. The risk of contamination and loss increases with increasing number of sample/flux transfers and thus, the GR method (three transfers) was deemed the most likely to lead to sample contamination and loss, resulting in errors in the analytical results. In addition, in this method, a brush is used to brush the mixture into the Pt–Au crucible, which needs to be cleaned to avoid contamination of the next sample. On the other hand, the ST method requires only one transfer, so that it has the lowest risk of contamination and loss. Contamination during the SH method (two transfers) can be avoided by using a small amount of deionized water, instead of a brush, to transfer any adherent sample. Thus, the risk of contamination and loss of the samples is of the order GR > SH > ST.

Based on our statistical calculations, the sample preparation time for the GR, ST, and SH methods were 6, 2, and 3 min, respectively. GR is the most time-consuming method but simultaneously provides information on the loss on ignition and major and minor elements from one sample. This method can save sample consumption and effectively avoids damage of the Pt–Au crucible by burning of the sample at high temperatures. This is because elements such as C and S are eliminated, while heavy metals such as Pb, Zn, Sn, and Sb are oxidized which, in the absence of burning, would be detrimental
to the Pt–Au crucible [30]. ST is the least time-consuming method; however, in this method, the Pt–Au crucible can be easily scratched by the glass rod during mixing. Finally, although slightly longer than that of ST, the SH sample preparation time is still acceptable. Notably, the elements in the rock samples are in oxide form (negligible poisoning), and the damage caused by the unburned samples to the Pt–Au crucible is therefore minimal. Thus, burning of the sample before fusing is not mandatory.

In summary, compared to the GR and ST methods, the SH method is a simpler and more effective sample preparation procedure and can be considered to be a trade-off procedure between uniformity, contamination and loss, and sample preparation time.

### 3.2. LLD

The LLD is defined as the lowest amount of analyte in the sample that can be detected but not necessarily quantitated under the stated experimental conditions [31]. The limit of quantitation (LOQ) is defined as the lowest concentration at which the analyte can be reliably detected and at which some predefined goals for bias and imprecision are met [32]. Notably, according to International Union of Pure and Applied Chemistry (IUPAC) guidelines, the LOQ is three-fold greater than the LLD [33]. In this study, the LLD was calculated from Equation (1) [34]:

\[
\text{LLD} = \frac{3}{s} \sqrt{\frac{r_{b}}{t_{b}}}
\]

where \(s\) is the sensitivity (cps/ppm), \(r_{b}\) is the estimated background intensity (cps) at the peak position, and \(t_{b}\) is the total background measurement time (s). The LLDs of the elements, automatically provided by SuperQ software, are listed in Table 2. Generally, the LLD is an important performance characteristic in method validation. The results indicate that the LLD of the fused glass discs produced by the three sample preparation procedures are all adequate for rock samples. No significant differences between the LLD of the SH and ST methods were observed. On the other hand, the LLD of MgO and CaO for these two methods were significantly higher than those of the GR method, while those of \(\text{Al}_2\text{O}_3\) and \(\text{Na}_2\text{O}\) were significantly lower.
Table 2. Data for the range of standard sample compositions and limits of detection (LLD) of the three sample preparation procedures.

| Number | Elements    | Range of Standard Sample Composition | LLD (µg g⁻¹) |
|--------|-------------|--------------------------------------|--------------|
| 1      | SiO₂        | 0.62–90.36                           | 157.94       |
| 2      | TiO₂        | 0.004–7.69                           | 25.73        |
| 3      | Al₂O₃       | 0.1–59.20                            | 271.70       |
| 4      | TFe₂O₃      | 0.075–25.65                          | 17.02        |
| 5      | MnO         | 0.001–0.43                           | 11.49        |
| 6      | MgO         | 0.006–49.40                          | 54.70        |
| 7      | CaO         | 0.04–51.10                           | 35.89        |
| 8      | Na₂O        | 0.008–10.59                          | 79.77        |
| 9      | K₂O         | 0.003–12.81                          | 13.82        |
| 10     | P₂O₅        | 0.002–6.06                           | 17.20        |
| 11     | Cr          | 1.6–15500                             | 11.13        |
| 12     | Cu          | 0.82–1230                            | 7.73         |
| 13     | Ba          | 6.4–4000                             | 53.28        |
| 14     | Ni          | 0.9–3780                             | 5.63         |
| 15     | Sr          | 2.3–12000                            | 3.82         |
| 16     | V           | 0.0022–768                           | 12.88        |
| 17     | Zr          | 0.7–1540                             | 2.06         |
| 18     | Zn          | 3.5–1300                             | 4.44         |

* m/m% is the mass percentage; TFe₂O₃ is the total iron oxide as ferric iron.

3.3. Accuracy and Precision

In this study, 19 CRMs were used to estimate the accuracy and precision of the three different sample preparation procedures for rock samples. The results, including the concentrations and standard deviations of ten major oxides and eight minor elements (WD-XRF), loss on ignition (LOI), and corresponding literature values are presented in Table S2. The sources of the recommend values of the 19 CRMs are summarized in Table S3.

In view of the numerous major oxide results (Table S2), which were not easy to compare, the data were sorted out and mapped as illustrated in Figures 4 and 5. Figure 4 presents the measured values of the major oxides obtained in this study and the recommended values cited from the literature. The solid line represents the regression of the measured results. The overall correlation coefficients of 10 major oxides are summarized in Table 3. This data indicated that the measured values of the different sample preparation procedures all show good correlation with the recommended values. However, it was difficult to determine which procedure was the best because the overall correlation coefficient values of the same element for the different procedures were very close. Thus, other criteria were required for comparison. Figure 5 displays the relative error values of the measured major oxide results. For SiO₂ and Al₂O₃, the relative error values were within 2%, except for RGM-2 which displayed relative errors in the range 2–3%. The points of relative error of SiO₂ and Al₂O₃ for the three sample preparation procedures were on the zero-point line, indicating good accuracy. For TFe₂O₃, MgO, CaO, K₂O, and NaO, the relative errors were within 5%, except for TFe₂O₃ in MGL-OshBO and JR-2 and MgO in JG-2, JR-2, and GH, which displayed very large relative errors; CaO in JR-2, AC-E, and GH with relative error values in the range 5–6%; and K₂O in JGb-2, PM-S, and AN-G with relative error values in the range 5–8%. This occurred because low element concentrations lead to small denominators and thus, a large relative error. The points of relative error of TFe₂O₃, CaO, K₂O, and NaO for the three sample preparation procedures did not differ much, indicating similar accuracies. One exception was MgO with points of relative error from the SH method being significantly closer to the zero-point line than those of the other two methods, thereby indicating better accuracy. In addition,
with increasing concentration, the points of relative error of TFeO$_2$, MgO, K$_2$O, and NaO for the three sample preparation procedures gradually converged to the zero-point line, indicating a gradual improvement in accuracy. Conversely, when the CaO concentration exceeded 12%, the points of relative error deviated from the zero-point line, probably due to the CaO calibration for downward deviation. For TiO$_2$, MnO, and P$_2$O$_5$, the relative errors were within 10%, except for TiO$_2$ in JG-2; MnO in JG-2 and RGM-2; and P$_2$O$_5$ in JG-2, AN-G, GH, JR-2, AC-E, JGb-2, W-2a, and MGL-OShBO. The points of relative error of TiO$_2$ and MnO significantly deviated from the zero-point line, indicating poor accuracy; this was attributed to a considerably low concentration. However, with the increase in the element concentration, the points of relative error of these two oxides approached the zero-point line, indicating that the accuracy improved with increasing concentration. Moreover, the TiO$_2$ from the SH method was significantly closer to the zero-point line, indicating superior accuracy. The points of relative error of P$_2$O$_5$ also deviated from the zero-point line when the concentration of P$_2$O$_5$ was <0.05%. Indeed, the concentration of P$_2$O$_5$ in JG-2 was lower than the LOQ, indicating that these data are incorrect. The concentrations of P$_2$O$_5$ in AN-G, GH, JR-2, AC-E, JGb-2, W-2a, and MGL-OShBO were too low so that the accuracy was poor. However, when the P$_2$O$_5$ concentration was >0.05%, the points were almost on the zero-point line and exhibited good accuracy.

![Figure 4. Representation of the measured and recommended values for the major oxides (letters (a), (b), and (c) represent the grinding (GR), stirring rod (ST), and shaker cup (SH) methods, respectively).](image)

**Table 3.** Results of the overall correlation coefficients between the measured and recommended values of the major oxides.

| Major Oxides | The Grinding (GR) Method | The Stirring Rod (ST) Method | The Shaker Cup (SH) Method |
|--------------|---------------------------|-----------------------------|---------------------------|
| SiO$_2$      | 0.99958                   | 0.99972                     | 0.99980                   |
| TiO$_2$      | 0.99710                   | 0.99893                     | 0.99939                   |
| Al$_2$O$_3$  | 0.99941                   | 0.99930                     | 0.99942                   |
| TFeO$_2$     | 0.99987                   | 0.99983                     | 0.99983                   |
| MnO          | 0.99961                   | 0.99649                     | 0.99664                   |
| MgO          | 0.99995                   | 0.99998                     | 0.99999                   |
| CaO          | 0.99967                   | 0.99960                     | 0.99954                   |
| Na$_2$O      | 0.99929                   | 0.99951                     | 0.99951                   |
| K$_2$O       | 0.99994                   | 0.99992                     | 0.99990                   |
| P$_2$O$_5$   | 0.99813                   | 0.99856                     | 0.99828                   |
were negative, indicating that the data were erroneous. Thus, the concentrations of Sr in JG-2 and GH, and Zn in JG-2 were higher than the LLD but lower than the LOQ, also indicating that the concentrations were inaccurate. In addition, the concentrations of Sr in JG-2 and MGL-OShBO were rather low and approached the LOQ, resulting in poor accuracy, while on the other hand, the analytical depth of Sr (3.8 mm) approached the thickness of the fused glass disc (3.6 mm).

The analytical depth of fluorescent X-rays is an extremely important parameter in XRF analysis since the fluorescent X-ray intensity increases with increasing thickness up to the analytical depth and remains constant thereafter [19]. Generally, the analytical depth of the specimen for each element can be calculated from its density, average elemental composition, and mass absorption coefficient, and the analytical depths of the fused glass disc for 18 elements calculated in this study are listed in Table 4. Considering that the thickness of a fused glass disc is 3.6 mm, it far exceeds the analytical depth for all major elements and most of the minor elements except for Sr (3.8 mm) and Zr (4.9 mm). Thus, for Sr and Zr, the thickness of the fused glass disc should be fixed to prevent variations in the fluorescent X-ray intensity. Notably, the error can be reduced by weighing constant sample and flux masses, maintaining constant conditions during sample preparation, and using the XRF instrument in rotation mode during measurement. However, the analytical depth of Sr approaches the thickness of the fused glass disc, in which a small change can lead to a significant change in the relative position between the thickness of the fused glass disc and analytical depth, thus affecting the accuracy and precision of the data.

For the minor elements (Table S2), the data attained were poor. The concentrations of Cr in JG-2, JR-2, RGM-2, AC-E, and GH; Cu in JG-2, JG-1a, JR-2, AC-E, GH, and MGL-OShBO; Ba in JGb-2, JR-2, GH, AN-G, and MGL-OShBO; Ni in JG-2, JR-2, RGM-2, AC-E, and GH; Sr in AC-E; and V in JG-2, JR-2, AC-E, GH, and MGL-OShBO were lower than the LLD, thereby indicating that the data were erroneous. Moreover, the concentrations of Cu in JH-1 and RGM-2, Ba in AC-E, Ni in JG-1a, Sr in JR-2 and GH, and Zn in JG-2 were higher than the LLD but lower than the LOQ, also indicating that the concentrations were inaccurate. In addition, the concentrations of Sr in JG-2 and MGL-OShBO were negative, indicating that the data were erroneous. Thus, the concentrations of Sr in JG-2 and MGL-OShBO were rather low and approached the LOQ, resulting in poor accuracy, while on the other hand, the analytical depth of Sr (3.8 mm) approached the thickness of the fused glass disc (3.6 mm).
Table 4. Analytical depths of the fluorescent X-rays in the fused glass disc.

| Analytical Line | Energy/keV | Analytical Depth/µm | Analytical Line | Energy/keV | Analytical Depth/µm |
|-----------------|------------|---------------------|-----------------|------------|---------------------|
| Si Kα           | 1.74       | 23                  | P Kα           | 2.01       | 21                  |
| Ti Kα           | 4.51       | 281                 | Cr Kα          | 5.41       | 530                 |
| Al Kα           | 1.49       | 10                  | Cu Kα          | 8.05       | 1080                |
| Fe Kα           | 6.40       | 560                 | Ba Lα          | 4.47       | 272                 |
| Mn Kα           | 5.90       | 461                 | Ni Kα          | 7.48       | 916                 |
| Mg Kα           | 1.25       | 4                   | Sr Kα          | 14.17      | 3831                |
| Ca Kα           | 3.69       | 158                 | V Kα           | 4.95       | 430                 |
| Na Kα           | 1.04       | 3                   | Zr Kα          | 15.78      | 4875                |
| K Kα            | 3.31       | 95                  | Zn Kα          | 8.64       | 1279                |

The afforded data revealed that relatively accurate results can be attained with the three sample preparation procedures for samples with concentrations higher than the LOQ. However, the vast data is overwhelming and difficult to compare. Therefore, it is necessary to quote a simple comparative mathematical analysis mode. Laurence Whitty-Léveillé et al. adopted a single value, which considered the accuracy and precision, to successfully compare three types of digestion methods and four types of analytical techniques [35]. Therefore, with reference to the Laurence Whitty-Léveillé method, we adopted two derivate equations from the residual sum of squares to compare the three sample preparation methods. These equations consider the standard deviations associated with the measured and recommended values but do not consider any possible systematic underestimation or overestimation by the different procedures due to the use of the squared error [35,36]:

\[ J_{S,j} = \frac{(x_{jitw} - x_{jlit})^2}{\sigma_{jitw}^2 + \sigma_{jlit}^2} \]  

\[ J_S = \sum_{i=1}^{N} J_{S,j} \]

For Equation (2), \( x_{jitw} \) is the measured value, \( x_{jlit} \) is the corresponding recommended literature value of \( x_{jitw} \), \( \sigma_{jitw} \) is the standard deviation, \( \sigma_{jlit} \) is the corresponding recommended literature value of \( \sigma_{jitw} \), and \( J_{S,j} \) is the dimensionless criterion value for the elements, which aids the comparison of elements with great absolute concentration variations. Based on Equation (2), accuracy is more important than precision. For Equation (3), \( N \) is the number of considered elements and \( J_S \) is the dimensionless criterion value for the samples. \( J_S \) provides an objective approach to estimate the best sample preparation procedure by summing the accuracy/precision of a given procedure in a single value (\( J_S \)). Considering the reproducibility of the measurement, the measured value better approaches the recommended value with decreasing \( J_S \).

Based on the data in Table S2, the \( J_{S,j} \) values were calculated and listed in Tables S4–S6. The \( J_S \) values of 57 CRMs with three sample preparation procedures were also calculated and listed in Table 5.

For the major oxides, the SH method presented the lowest \( J_{S,\text{major}} \) value (3.95), followed by those attained from the GR (5.57) and ST (5.94) methods. On the other hand, for the minor elements, the ST method presented the lowest \( J_{S,\text{minor}} \) value (11.36), followed by those attained by the SH (16.64) and GR (33.09) methods. Based on the work of Whitty-Léveillé, the measured values better approach the recommended values with decreasing \( J_{S,\text{major}} \) and \( J_{S,\text{minor}} \) values. These data indicate that the GR method is a poor choice for the major oxides and minor elements. If both the major oxides and minor elements are considered, then the ST method displays a lower \( J_S \) value (16.93) than that of the SH method (20.59). However, notably, the biggest difference between the two methods is from Zr, with a \( J_{S,\text{without Zr}} \) value of 0.87 for the ST method and 5.39 for the SH method. Thus, if Zr is excluded, the SH method shows a lower \( J_{S,\text{without Zr}} \) value (15.20) than that of the ST method (16.06). We therefore recommend the use of the \( J_{S,\text{without Zr}} \) value to evaluate the three methods, whereby the SH method...
shows the lowest value indicating the best accuracy and precision. As for Zr, the reason for the poor data result by the SH method is yet unknown. We intend to investigate this in the future.

Table 5. Weighted discrepancies between the measured and recommended values of 19 reference materials obtained by the three different sample preparation procedures and analyzed by wavelength dispersive X-ray fluorescence spectrometry (WD-XRF).

| Elements | JST,i a | n b | JSA,i c | JSW,i d |
|----------|---------|-----|---------|---------|
| Major    |         |     |         |         |
| SiO₂     | 5.54    | 5.47| 4.28    | 17      | 0.33 | 0.32 | 0.25 |
| TiO₂     | 16.96   | 7.2 | 4.56    | 16      | 1.06 | 0.45 | 0.29 |
| Al₂O₃    | 5.44    | 6.39| 3.27    | 17      | 0.32 | 0.38 | 0.19 |
| TFe₂O₃   | 2.03    | 16.06| 11.6   | 17      | 0.12 | 0.94 | 0.68 |
| MnO      | 30.57   | 5.62| 5.97    | 17      | 1.80 | 0.33 | 0.35 |
| MgO      | 9.62    | 8.1 | 5.3     | 16      | 0.60 | 0.51 | 0.33 |
| CaO      | 16.05   | 23.96| 21.59 | 17      | 0.94 | 1.41 | 1.27 |
| Na₂O     | 1.33    | 1.46| 2.53    | 17      | 0.08 | 0.09 | 0.15 |
| K₂O      | 2.91    | 7.32| 2.91    | 17      | 0.17 | 0.43 | 0.17 |
| P₂O₅     | 8.31    | 11.43| 4.25   | 16      | 0.52 | 0.71 | 0.27 |
| Minor    |         |     |         |         |
| Cr       | 9.65    | 3.71| 4       | 12      | 0.80 | 0.31 | 0.33 |
| Cu       | 87.84   | 22.48| 24.1   | 9       | 9.76 | 2.50 | 2.68 |
| Ba       | 2.57    | 5.24| 7.71    | 12      | 0.21 | 0.44 | 0.64 |
| Ni       | 11.35   | 7.72| 5.57    | 10      | 1.14 | 0.77 | 0.56 |
| Sr       | 55.94   | 40.44| 44.52  | 12      | 4.66 | 3.37 | 3.71 |
| V        | 7.6     | 3.61| 4.96    | 11      | 0.69 | 0.33 | 0.45 |
| Zr       | 14      | 14.83| 91.66  | 17      | 0.82 | 0.87 | 5.39 |
| Zn       | 240.05  | 44.41| 45.96  | 16      | 15.00| 2.78 | 2.87 |
| JST, major d | 5.94 | 5.57| 3.95    | 17      | 0.33 | 0.38 | 0.19 |
| JST, minor e | 33.09| 11.36| 16.64 | 17      | 0.33 | 0.38 | 0.19 |
| JST, f    | 39.03   | 16.93| 20.59  | 16      | 0.33 | 0.38 | 0.19 |
| JST, without Zr g | 38.21| 16.06| 15.20 | 16      | 0.33 | 0.38 | 0.19 |

a JST,i is the sum of the JSi values of an element from different samples; b n is the total number of the effective measured value of an element; c JSA,i is the value of JST,i divided by n; d JST, major is the sum of the JSA,i values of major oxides; e JST, minor is the sum of the JSA,i values of minor elements; f JST is the sum of the JSA,i values of major oxides and minor elements; g JST,without Zr is the sum of the JSA,i values of elements including SiO₂, TiO₂, Al₂O₃, TFe₂O₃, MnO, MgO, CaO, Na₂O, K₂O, P₂O₅, Cr, Cu, Ba, Ni, Sr, V, and Zn.

In this paper, the values of the LOI of 19 CRM s are also listed in Table S2. For JG-2, John Stix et al. have reported a value of 0.41 [37], while the value reported by Antje Herbrich et al. and Roman Golowin et al. was 0.57 [38,39]. The LOI of JG-2 in this study is the same as that reported by John Stix. For JG-1a, John Stix et al. have reported a value of 1.21 [37], while Zongshou Yu et al. have reported a value of 0.55 [40]. The LOI value of JG-1a in this study was 0.68, which approaches that reported by Zongshou Yu. For JA-3, Antje Herbrich et al. and Roman Golowin et al. have reported a value of 0.34 [38,39], John Stix et al. reported a value of 1.08 [37], and Sarah Freund et al. a value of 0.12 [41]. The LOI value of JA-3 in this study was 0.18, which is similar to that reported by Sarah Freund. For JR-2, the LOI value in this study is consistent with that reported by John Stix [37]. For GSP-2, Ingrid Raczek et al. reported a value of 0.96 [42], Zongshou Yu et al. a value of 0.8 [40], and Lindsay J. McHenry a value of 2.88 [43]. The LOI value of GSP-2 in this study was 1.12, which is close to that reported by Ingrid Raczek. For BCR-2, many different values (−0.04, −0.02, −0.01, 0, 0.07, and 0.69) were measured by other authors [40, 42, 44–49], while the LOI value in this study was −0.06. For BHVO-2, many different values ranging from −0.88 to 0.97 were measured by other authors [40, 42–47, 49–55], while the LOI in this study was −0.55. For JH-1, the LOI in this study was 2.13, which is higher than the recommended value of 1.78. On the other hand, for JGb-2, AC-E, GH, PM-S, WS-E, AN-G, and MGL-OShBO, the LOI
values in this study well agree with the literature recommended values. Finally, the JB-2a, JB-3a, W-2, and RGM-2 LOI values are reported for the first time herein.

4. Conclusions

In this study, the advantages and disadvantages of three fusion technique sample preparation procedures for the routine XRF analysis of vast rock samples were compared. In terms of the operational process, the SH method greatly reduced the sample preparation time compared to that of the GR method. It also afforded a more uniform mixture prior to fusing over that attained with the ST method. In terms of the detection limits, the LLD of MgO and CaO attained by the SH method were significantly higher than those attained by the GR method, while those of Al$_2$O$_3$ and Na$_2$O were significantly lower. The LLD of the other tested elements were near identical for the three methods. Additionally, we adopted a dimensionless criterion index, $J_{S;i}$ without Zr, to evaluate the accuracy and precision. The $J_{S;i}$ without Zr value of the SH method was the lowest when Zr was excluded, suggesting that of the three tested methods, this method afforded the most accurate and precise data. Overall, the SH method is highly recommended as a better sample preparation procedure over the other two tested methods.

Supplementary Materials: The following are available online at http://www.mdpi.com/1424-8220/20/18/5325/s1, Figure S1: Fused glass disc preparation procedure, Figure S2: Procedures for mixing the samples and flux by (a) grinding, (b) stirring, and (c) shaking, Table S1: Details of the certified reference materials (CRMs), Table S2: Results of the major and minor element determination by X-ray fluorescence (XRF) analysis of the samples prepared by the three preparation procedures, Table S3: Source of recommend values of 19 CRMs, Table S4: The values of $J_{S;i}$ by grinding method, Table S5: The values of $J_{S;i}$ by stirring rod method, Table S6: The values of $J_{S;i}$ by shaker cup method.

Author Contributions: D.-P.Z., D.-S.X. and Y.-H.L. made contributions to design the methods. D.-P.Z., Q.G. and J.-J.G. made contributions to the experiment and data collection. D.-P.Z., D.-S.X. and B.W. took part in manuscript preparation and editing. All authors have read and agreed to the published version of the manuscript.

Funding: This paper was financially supported by the National Key R&D Program of China (2019YFA0708400); Experimental Technology Innovation Fund of the Institute of Geology and Geophysics, Chinese Academy of Sciences (Grant no. T202001); Equipment function development technology innovation projects of China science academy (Grant no. IGG201702); National Natural Science Foundation of China (Grant nos. 41403021, 41703021, and 41525012); and Open Research Foundation of the Key Laboratory of Mineral Resources, Institute of Geology and Geophysics, Chinese Academy of Sciences (Grant no. 12).

Conflicts of Interest: The authors declare no conflict of interest.

References

1. Li, X.; Zou, H.; Zhang, Q.; Wei, S.; Wang, L.; Zhang, Y.; Yang, J. Application and analysis of portable X-ray fluorescence analyzer on fluorite exploration in shallow cover area. Comput. Tech. Geophys. Geochem. Explor. 2018, 40, 681–688.
2. Meng, L.; Han, P.; Wang, S.; Ren, D.; Wang, J. Application progress of X-ray fluorescence spectroscopy in the detection of heavy metals in soil. Food Mach. 2017, 33, 210–213.
3. Bounakha, M.; Ernbarch, K.; Zahry, F.; Biai, E.; Kump, P. Capabilities of elemental analysis by EDXRF for geochemistry. J. Radioanal. Nucl. Chem. 2008, 275, 467–478. [CrossRef]
4. Leoni, L.; Menichini, M.; Saitta, M. Determination of S, Cl and F in silicate rocks by X-Ray fluorescence analyses. X-ray Spectrom. 1982, 11, 156–158. [CrossRef]
5. Ren, J.; Yan, R.; Liu, Y. Determination of trace chromium in cement by X-ray fluorescence spectrometer. Chin. J. Anal. Lab. 2015, 34, 735–739.
6. Xie, L.; Wang, J.; Zhao, X.; Yang, R.; Feng, X.; Li, Y.; Luan, L.; Zhang, L.; He, R.; Liu, Y.; et al. Rapid screening of chromium-tainted gelatin capsules by wavelength dispersive X-ray fluorescence spectrometry. Chin. J. Pharm. Anal. 2013, 33, 286–288, 291.
7. Smagunova, A.N.; Rozova, O.F.; Skribko, N.N. X-ray fluorescence analysis of powder products in ferrous-metallurgy-review. Ind. Lab. 1990, 56, 1062–1073.
8. Pfandl, K.; Kueppers, B.; Scheiber, S.; Stockinger, G.; Holzer, J.; Pomberger, R.; Antrekowitsch, H.; Vollprecht, D. X-ray fluorescence sorting of non-ferrous metal fractions from municipal solid waste incineration bottom ash processing depending on particle surface properties. Waste Manag. Res. 2019. [CrossRef] [PubMed]
9. Leake, B.E.; Hendry, G.L.; Kemp, A.; Plant, A.G.; Harrey, P.K.; Wilson, J.R.; Coats, J.S.; Aucott, J.W.; Lünel, T.; Howarth, R.J. The chemical analysis of rock powders by automatic X-ray fluorescence. Chem. Geol. 1969, 5, 7–86. [CrossRef]
10. Afonin, V.P. X-ray fluorescence analysis of rocks. Fresenius’ Z. Anal. Chem. 1989, 335, 54–57. [CrossRef]
11. Nakayama, K.; Nakamura, T. X-ray fluorescence analysis of rare earth elements in rocks using low dilution glass beads. Anal. Sci. 2005, 21, 815–822. [CrossRef] [PubMed]
12. Norrish, K.; Hutton, J.T. An accurate X-ray spectrometric method for the analysis of a wide range of geological samples. Geochim. Cosmochim. Acta 1969, 33, 431–453. [CrossRef]
13. Lodha, G.S.; Sawhney, K.J.S.; Choubey, V.M. Determination of Y, Zr, Ba, La and Ce in granitic rocks using energy-dispersive X-ray fluorescence. X-ray Spectrom. 1989, 18, 225–227. [CrossRef]
14. Eastell, J.; Willis, J.P. A low dilution fusion technique for the analysis of geological samples I—Method and trace element analysis. X-ray Spectrom. 1990, 19, 3–14. [CrossRef]
15. Eastell, J.; Willis, J.P. A low dilution fusion technique for the analysis of geological samples. 2—Major and minor element analysis and the use of influence/alpha coefficients. X-ray Spectrom. 1993, 22, 71–79. [CrossRef]
16. Goto, A.; Tatsumi, Y. Quantitative analysis of rock samples by an X-ray fluorescence spectrometer (I). Rigaku J. 1994, 11, 40–59.
17. Goto, A.; Tatsumi, Y. Quantitative analysis of rock samples by an X-ray fluorescence spectrometer (II). Rigaku J. 1996, 13, 20–38.
18. Tamponi, M.; Bertoli, F.; Innocenti, F.; Leoni, L. X-ray fluorescence analysis of major elements in silicate rocks using fused glass discs. Atti Soc. Tosc. Sci. Nat. Mem. 2003, 108, 73–79.
19. Ichikawa, S.; Nakamura, T. Solid Sample Preparations and Applications for X-Ray Fluorescence Analysis. In Encyclopedia of Analytical Chemistry; John Wiley & Sons, Ltd.: Chichester, UK, 2016; pp. 1–22. [CrossRef]
20. Ogasawara, M.; Mikoshiba, M.; Geshi, N.; Shimoda, G.; Ishizuka, Y. Optimization of analytical conditions for major element analysis of geological samples with XRF using glass beads. Bull. Geol. Surv. Jpn. 2018, 69, 91–103. [CrossRef]
21. Tani, K.; Kawabata, H.; Chang, Q.; Sato, K.; Tatsumi, Y. Quantitative analyses of silicate rock major and trace elements by X-ray fluorescence spectrometer: Evaluation of analytical precision and sample preparation. Front. Res. Earth Evol. 2005, 2, 1–8.
22. Orihashi, Y.; Hirata, T. Rapid quantitative analysis of Y and REE abundances in XRF glass bead for selected GSJ reference rock standards using Nd-YAG 266 nm UV laser ablation ICP-MS. Geochem. J. 2003, 37, 401–412. [CrossRef]
23. Stern, W.B. On trace element analysis of geological samples by X-ray fluorescence. X-ray Spectrom. 1976, 5, 56–60. [CrossRef]
24. Yamasaki, T. XRF major element analyses of silicate rocks using 1:10 dilution ratio glass bead and a synthetically extended calibration curve method. Bull. Geol. Surv. Jpn. 2014, 65, 97–103. [CrossRef]
25. Kamei, A. Determination of trace element abundances in GSJ reference rock samples using lithium metaborate–lithium tetraborate fused solutions and inductively coupled plasma mass spectrometry. Geosci. Rept. Shimane Univ. 2016, 34, 41–49.
26. Krusberski, N. Exploring Potential Errors in XRF Analysis; The Southern African Institute of Mining and Metallurgy Analytical Challenges in Metallurgy: Johannesburg, South Africa, 2006; pp. 1–8.
27. GmbH, R. Representative Sample Preparation for XRF Analysis. AZoM. Available online: https://www.azom.com/article.aspx?ArticleID=16242 (accessed on 8 September 2020).
28. Muller, R. Spectrochmical Analysis by X-ray Fluorescence; Plenum Press: New York, NY, USA, 1972.
29. Anzelmo, J.; Bouchard, M.; Provencher, M.E. Atomic Perspectives X-ray Fluorescence Spectroscopy, Part II: Sample Preparation. Spectrosc. Springf. Eugene Duluth 2014, 29, 12.
30. Lupton, D.F.; Merker, J.; Schötz, F. The Correct Use of Platinum in the XRF Laboratory. X-ray Spectrom. 1997, 26, 132–140. [CrossRef]
31. Shrivastava, A.; Gupta, V. Methods for the determination of limit of detection and limit of quantitation of the analytical methods. Chron. Young Sci. 2011, 2, 21–25. [CrossRef]
32. Armbruster, D.A.; Pry, T. Limit of blank, limit of detection and limit of quantitation. *Clin. Biochem. Rev.* 2008, 29 (Suppl. 1), S49–S52.

33. Gazulla, M.F.; Rodrigo, M.; Vicente, S.; Orduña, M. Methodology for the determination of minor and trace elements in petroleum cokes by wavelength-dispersive X-ray fluorescence (WD-XRF). *X-ray Spectrom.* 2010, 39, 321–327. [CrossRef]

34. Han, X. Determination of Palladium, Platinum and Gold by X-ray Fluorescence Spectrometry with Resin Separation; Yantai University: Yantai, China, 2013.

35. Whitty-Leveil, L.; Turgeon, K.; Bazin, C.; Larivière, D. A comparative study of sample dissolution techniques and plasma-based instruments for the precise and accurate quantification of REEs in mineral matrices. *Anal. Chim. Acta* 2017, 961, 33–41. [CrossRef]

36. Anscoumbe, F.J.; Tukey, J.W. The Examination and Analysis of Residuals. *Technometrics* 1963, 5, 141–160. [CrossRef]

37. Stix, J.; Gorton, M.P.; Fontaine, E. Major and trace element analysis of fifteen Japanese igneous reference rocks by XRFs and INAA. *Geostand. News.* 1996, 20, 87–94. [CrossRef]

38. Herbrich, A.; Hauff, F.; Hoernle, K.; Werner, R.; Garbe-Schönberg, D.; White, S. A 1.5 Ma record of plume-ridge interaction at the Western Galápagos Spreading Center (91°40′–92°00′ W). *Geochim. Cosmochim. Acta* 2016, 185, 141–159. [CrossRef]

39. Golowin, R.; Portnyagin, M.; Hoernle, K.; Hauff, F.; Werner, R.; Garbe-Schönberg, D. Geochemistry of deep Manihiki Plateau crust: Implications for compositional diversity of large igneous provinces in the Western Pacific and their genetic link. *Chem. Geol.* 2018, 493, 553–566. [CrossRef]

40. Yu, Z.; Norman, M.D.; Robinson, P. Major and trace element analysis of silicate rocks by XRF and laser ablation ICP-MS using lithium borate fused glasses: Matrix Effects, Instrument Response and Results for International Reference Materials. *Geostand. News.* 2003, 27, 67–89. [CrossRef]

41. Freund, S.; Haase, K.M.; Keith, M.; Beier, C.; Garbe-Schönberg, D. Constraints on the formation of geochemically variable plagiogranite intrusions in the Troodos Ophiolite, Cyprus. *Contrib. Mineral. Petrol.* 2014, 167, 978. [CrossRef]

42. Raczek, I.; Stoll, B.; Hofmann, A.W.; Peter Jochum, K. High-precision trace element data for the USGS reference materials BCR-1, BCR-2, BHVO-1, BHVO-2, AGV-1, AGV-2, DTS-1, DTS-2, GSP-1 and GSP-2 by ID-TIMS and MIC-SSMS. *Geostand. News.* 2001, 25, 77–86. [CrossRef]

43. McHenry, L.J. Element mobility during zeolitic and argillic alteration of volcanic ash in a closed-basin lacustrine environment: Case study Olduvai Gorge, Tanzania. *Chem. Geol.* 2009, 265, 540–552. [CrossRef]

44. Zhang, G.; Smith-Duque, C.; Tang, S.; Li, H.; Zarikian, C.; D’Hondt, S.; Inagaki, F. Geochemistry of basalts from IODP site U1365: Implications for magmatism and mantle source signatures of the mid-Cretaceous Osburn Trough. *Lithos* 2012, 144–145, 73–87. [CrossRef]

45. Zhang, G.-L.; Chen, L.-H.; Li, S.-Z. Mantle dynamics and generation of a geochemical mantle boundary along the East Pacific Rise–Pacific/Antarctic ridge. *Earth Planet. Sci. Lett.* 2013, 383, 153–163. [CrossRef]

46. Taneja, R.; Rushmer, T.; Blichert-Toft, J.; Turner, S.; O’Neill, C. Mantle heterogeneities beneath the Northeast Indian Ocean as sampled by intra-plate volcanism at Christmas Island. *Lithos* 2016, 262, 561–575. [CrossRef]

47. Allaway, B.V.; Pearce, N.J.G.; Villarosa, G.; Outes, V.; Moreno, P.I. Multiple melt bodies fed the AD 2011 eruption of Puyehue-Cordillera Caulle, Chile. *Sci. Rep.* 2015, 5, 17589. [CrossRef] [PubMed]

48. Jung, S.; Viyet, K.; Romer, R.L.; Mezger, K.; Hoernes, S.; Satir, M. Petrogenesis of Tertiary Alkaline Magmas from IODP site U1365: Implications for magmatism and mantle source signatures of the mid-Cretaceous Osburn Trough. *Lithos* 2012, 144–145, 73–87. [CrossRef]

49. Wei, F.; Prytulak, J.; Xu, J.; Wei, W.; Hammond, J.O.S.; Zhao, B. The cause and source of melting for the most recent volcanism in Tibet: A combined geochemical and geophysical perspective. *Lithos* 2017, 288–289, 175–190. [CrossRef]

50. Guo, J.; Guo, F.; Yan Wang, C.; Li, C. Crustal recycling processes in generating the early Cretaceous Fangcheng basalts, North China Craton: New constraints from mineral chemistry, oxygen isotopes of olivine and whole-rock geochemistry. *Lithos* 2013, 170–171, 1–16. [CrossRef]

51. Cheng, C.; Zheng, H.; Kapsiotis, A.; Liu, W.; Lenaz, D.; Velicogna, M.; Zhong, L.; Huang, Q.; Yuan, Y.; Xia, B. Geochemistry and geochronology of dolerite dykes from the Daba and Dongbo peridotite massifs, SW Tibet: Insights into the style of mantle melting at the onset of Neo-Tethyan subduction. *Lithos* 2018, 322, 281–295. [CrossRef]
52. Phillips, E.H.; Sims, K.W.W.; Sherrod, D.R.; Salters, V.J.M.; Blusztajn, J.; Dulai, H. Isotopic constraints on the genesis and evolution of basanitic lavas at Haleakala, Island of Maui, Hawaii. *Geochim. Cosmochim. Acta* 2016, 195, 201–225. [CrossRef]

53. Gao, R.; Lassiter, J.C.; Ramirez, G. Origin of temporal compositional trends in monogenetic vent eruptions: Insights from the crystal cargo in the Papoose Canyon sequence, Big Pine Volcanic Field, CA. *Earth Planet. Sci. Lett.* 2017, 457, 227–237. [CrossRef]

54. Liu, H.-Q.; Xu, Y.-G.; Tian, W.; Zhong, Y.-T.; Mundil, R.; Li, X.-H.; Yang, Y.-H.; Luo, Z.-Y.; Shang-Guan, S.-M. Origin of two types of rhyolites in the Tarim Large Igneous Province: Consequences of incubation and melting of a mantle plume. *Lithos* 2014, 204, 59–72. [CrossRef]

55. He, B.; Zhong, Y.-T.; Xu, Y.-G.; Li, X.-H. Triggers of Permo-Triassic boundary mass extinction in South China: The Siberian Traps or Paleo-Tethys ignimbrite flare-up? *Lithos* 2014, 204, 258–267. [CrossRef]

© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).