Mineralogical and Geochemical Study on the Yaojiazhuang Ultrapotassic Complex, North China Craton: Constraints on the Magmatic Differentiation Processes and Genesis of Apatite Ores

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The differentiation process of ultrapotassic magmas is enigmatic and poorly understood. The Yaojiazhuang ultrapotassic complex is concentrically zoned by late-intruded syenite in the core and early emplaced clinopyroxenite in the periphery, combining a “bi-modal” lithology. Spatially, apatite and iron oxide-apatite (IOA) ores, glimmerite and pseudoleucite occur in the upper part of clinopyroxenite. The syenite and clinopyroxenite are composed of variable amounts of clinopyroxene, biotite, K-feldspar, magnetite, apatite with minor analcite, titanite, and primary calcite. The pseudoleucite clinopyroxenite contains mainly clinopyroxene, biotite and garnet in the matrix, and nepheline–K-feldspar intergrowth with muscovite and minor celestine in the leucite pseudomorph. Geochemically, rocks of the Yaojiazhuang complex are significantly enriched in potassium (K), light rare earth elements (LREE), and large ion lithophile elements (LILE). Crustal contamination by Archean tonalite–trondhjemite–granodiorite (TTG) gneisses basement may play an important role to convert the syenitic melts from silica-undersaturation to saturation. Fractionation crystallization is supported by the mineral crystallization sequence to explain the bimodal lithologies instead of silicate liquid immiscibility. During the magmatic evolution, decompression, fractionation of volatile-poor clinopyroxene and the enhancement by CO₂ may result in the exsolution of an aqueous fluid. The late-stage interactions between existing minerals and magmatic fluids in the crystal mush could be a key process in the generation of both leucite pseudomorphs and apatite/IOA ores.

Keywords: ultrapotassic, Yaojiazhuang complex, fluid exsolution, apatite, pseudoleucite, syenite, clinopyroxenite, North China Craton

INTRODUCTION

The occurrence of ultrapotassic rocks is volumetrically rare, with the origin usually related to intraplate extension environments and collision zones (e.g., Foley et al., 1987; Miller et al., 1999; Conticelli et al., 2007, 2009, 2013). Considerable amounts of studies had been conducted on ultrapotassic rocks which are mostly focusing on their mantle source and partial melting.
processes (e.g., Thompson et al., 1990; Conticelli and Peccherillo, 1992; Foley and Peccherillo, 1992; Rogers et al., 1992; Zhang et al., 1995; Miller et al., 1999; Lobach-Zhuchenko et al., 2008; Gaeta et al., 2016). Metasomatized mantle sources and relatively low degree of melting process had been proposed for the generation of ultrapotassic primary melts (e.g., Dawson, 1987; Marks et al., 2008; Sun et al., 2014; Bodeving et al., 2017; Soder and Romer, 2018; Soköl et al., 2018). Nevertheless, most of the ultrapotassic rocks are not solidified from primary mantle-derived liquids, thus the subsequent differentiation processes of ultrapotassic magmas in the crust including fractionation crystallization, crustal contamination, magma (un-)mixing and mingling, and exsolution/transportation of both magmatic and post-magmatic fluids could also play important roles in the formation of them (e.g., Ferlito and Lanzafame, 2010; Jeffery et al., 2013; Brenna et al., 2015; Wolff, 2017; Burchardt, 2018). However, to the best of our knowledge, these crustal processes are poorly constrained. Hence, a comprehensive illustration on the differentiation processes is crucial for our understanding about the formation of ultrapotassic rocks.

Moreover, some alkaline rocks, including ultrapotassic rocks are associated with mineral deposits, mainly apatite ± magnetite ± minor silicate minerals (phoscorite, IOA ores and monomineralic apatite ores), such as Ordovician-Silurian Misvaerdal complex in Norway (Ihlen et al., 2014), Khibiny complex in Kola Peninsula, Russia (Notholt, 1979; Kogarko, 2018) and Mushgai-Khudag Complex, South Mongolia (Nikolenko et al., 2018). Accordingly, the formation of these P-rich ores is genetically related to the differentiation of the alkaline magmas. Two major models including fractional crystallization (Hou et al., 2015) and liquid immiscibility (Kolker, 1982; Jiang et al., 2004; Mokhtari et al., 2013) had been proposed for the formation of these apatite and IOA ores. Therefore, it is of great significance to elucidate the petrogenesis of ore-bearing alkaline complex, which could potentially shed new lights on the formation of the apatite and IOA ores.

The Yaojiazhuang alkaline plutonic intrusion is a "bi-modal" ring complex with syenite in the core and clinopyroxenite in the periphery generated by ultrapotassic magmas frozen in the crust according to previous studies (Chen et al., 2013). Notably, there is also a large apatite and IOA ore deposit hosted at the upper part of clinopyroxenite and have been mined as phosphorus deposit for decades. However, the formation mechanism of bi-modal lithofacies and tons of phosphorus concentration are still in widely debates and remain poorly understood (e.g., Hou, 1990b; Mu and Yan, 1992; Zhang, 1999; Chen et al., 2013; Tang et al., 2014; Liu and Tang, 2018). Previous studies suggested that the compositionally contrasting syenite and clinopyroxenite could be the Si-rich and Fe-rich melts produced by silicate liquid immiscibility from a common ultrapotassic parental magma, supported by thermodynamic simulations (Hou, 1990b; Ma et al., 1999), and during the separation of immiscible liquids, P_2O_5 tends to be partitioned into the Fe-rich melts and thus form the apatite and IOA ore in clinopyroxenite (Hou, 1990a). Alternatively, other authors commonly favor fractional crystallization either for Yaojiazhuang complex (Chen et al., 2013) or adjacent Fanshan complex (Cheng and Sun, 2003; Hou et al., 2015).

Additionally, the widespread hydrated minerals like biotite and the occurrence of pegmatites which has not been reported in detail indicate that during the formation of Yaojiazhuang complex, the magma ought to be enriched in fluids. Recent studies had revealed that the interactions between fluids and minerals during the last magmatic stage could significantly modify rock structures and textures, and could also act as a key mechanism in the generation of economic ore deposits, especially apatite and IOA deposits (e.g., Roedder, 1992; Huber et al., 2012; Ballhaus et al., 2015; Guo and Audet, 2017; La Cruz et al., 2019; Knipping et al., 2019). Therefore, a combined study involving mineralogy, petrology and geochemistry studies is required to elucidate these processes. Thus in this paper, we conducted comprehensive study for the differentiation of Yaojiazhuang complex, in order to better understand the differentiation processes of ultrapotassic magmas and shed new lights on the formation of the apatite and IOA ores.

GEOLoGICAL SETTING

Regional Geology

The Yaojiazhuang ultrapotassic complex is located at the north margin of the North China Craton (NCC), ~160 km northwest of the Beijing city (Figure 1A). The basement rocks of NCC are mainly mafic granulites, amphibolites and tonalite-trondhjemite–granodiorite (TTG) gneisses of ages up to 3.85 Ga (Liu et al., 1992), and are overain unconformably by Mesozoic–Neo-Proterozoic sediments (Zhao et al., 2001). To the north of the NCC is the Central Asian Orogenic Belt, which strongly influenced the northern margin of NCC during Carboniferous to Permian by the progressive closure of Paleo-Asian Ocean leading to Andean-type magmatism and Himalayan-type collision, and the formation of the Solonker suture (e.g., Xiao et al., 2003; Zhang et al., 2009). Following the closure was a post-collisional extension during early Mesozoic, resulting in a series of magmatic emplacement events and forming a > 1,500 km E-W trending alkaline rock belt along the northern margin of NCC (e.g., Zhang and Wang, 1997; Zhang, 1999; Zhao et al., 2008; Ren et al., 2009; Zhang et al., 2012). The major rock types can be divided mainly into three associations: (1) clinopyroxene diorite – quartz syenite – aegirine syenite – syenite – granite association, such as Shuiqangou complex; (2) nepheline syenite – aegirine syenite – urtite – jilolite association, such as Xiangshuigou and Aoyukou complexes; (3) clinopyroxene – biotite clinopyroxene – nelsonite – syenite association, representative complexes usually host economic iron and phosphorus ore deposits such as Yaojiazhuang (this study) and Fanshan complexes (e.g., Mu and Yan, 1992; Zhang, 1999; Jiang et al., 2004; Zhang et al., 2012; Hou et al., 2015).

Geology of Yaojiazhuang Ultrapotassic Complex

The ca. 221 Ma Yaojiazhuang ultrapotassic complex (Zhang et al., 2012) is an oval-shaped intrusive body in plan with a
FIGURE 1 | (A) Simplified tectonic map showing the distribution of Late Triassic alkaline intrusions in North China Craton (modified from Ren et al., 2009). Labeled ages and analytical methods are compiled from Zhang et al. (2012) and references therein. (B) Geological map of Yaojiazhuang ultrapotassic complex modified from Hou (1990a).
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size around 1.8 × 1.3 km (Figure 1B). It intruded the Archean gneiss and consists of a “bi-model” lithology with syenite at the core and clinopyroxenite at the periphery. At the places where they directly contacted, mingling between syenite and clinopyroxenite is observed (Figure 2A). In some parts of the complex, K-feldspar clinopyroxenite is outcropped either between the units of syenite and clinopyroxenite or scattered adjacent to the clinopyroxenite (Figure 2B). Besides, some clinopyroxenite host cm-sized pseudoleucite which had been previously identified as K-feldspar clinopyroxenite. At the upper part of clinopyroxenite, i.e., toward to syenite, the amount of apatite increased continuously, forming porphyritic or spheroidal apatite + magnetite ores up to around 10 cm (Figure 2C). These apatite clusters are cemented by a mineral assemblage resemble to clinopyroxenite. Coarsening of biotite is also observed locally, thus rocks in these places become extremely fragile (Figure 2D). Pegmatitic syenites occurred at the center of the syenite unit (Figure 2E) and aegirine syenite is locally present. Moreover, except the main units as mentioned above, late stage intrusive activities are quite vigorous, almost

FIGURE 2 | Field photos showing the geology of Yaojiazhuang ultrapotassic intrusion. (A) Angular clinopyroxenite breccia at the boundary between syenite and clinopyroxenite. (B) Magma mingling between syenite and clinopyroxenite. (C) Intrusive contact between clinopyroxenite and K-feldspar clinopyroxenite. (D) Spheroidal apatite ores in clinopyroxenite. (E) Pegmatitic syenite at the center of the syenite unit. (F) Clinopyroxenite, syenite and pegmatitic syenite feeder dykes in the clinopyroxenite.
all the rocks in the Yaojiazhuang complex are cut by dykes with various compositions (Figure 2F).

**ANALYTICAL METHOD**

**Scanning Electron Microscope**

Polished thin sections were firstly mapped using a Zeiss Ultra 55 FESEM equipped with an INCA MAX 20 electron dispersive spectrometry (EDS) system at the Key Laboratory of Submarine Geosciences State Oceanic Administration, Second Institute of Oceanography, Ministry of Natural Resources (SIOMNR), China. The employed accelerating voltage was 15 kV.

**Whole-Rock Major and Trace Elements**

Samples were firstly peeled off the weathered or altered surfaces before jaw-crushing and the subsequent grounding in agate mills to powders of over 200 mesh. The pseudoleucite clinopyroxenites were initially packaged in a plastic envelope and crushed by hammering to small pieces, and then those pieces of as much as pure pseudoleucite or matrix were separately collected by hands and grounded to powders of ~200 mesh in an agate mill. Major elements were analyzed by X-ray fluorescence (XRF) on fused glass discs at the laboratory of Nanjing Hongchuang Exploration Technology Service Co., Ltd. (NHETS) in Nanjing, China. The analytical uncertainties are < 1%, estimated from repeated analyses of 10 standards (GBW07101, GBW07103, GBW07105, GBW07114, AGV-2, BHVO-2, COQ-1, DTS-2b, GSP-2, and W-2a). Trace element data was determined by inductively-coupled–plasma mass-spectrometry (ICP-MS) after acid digestion of powders in high-pressure Teflon bullets performed at NHETS. The analytical precision for trace and rare earth elements is generally better than 10%, verified by repeated analyses of eight standards (GBW07101, GBW07316, BCR-2, W-2a, GSP-2, DTS-2a, BHVO-2, and AGV-2).

**Electron Microprobe Analysis**

Compositions of mineral phases were analyzed at SIOMNR using a JXA-8100 electron probe micro-analyzer (EPMA). Operating conditions were performed at 15 kV accelerating voltage with a 20 nA beam current. The beam size was 5 µm or 1 and up to 15 µm depending on crystal sizes. The peaks are counted by 10 s with background half of that for every element and ZAF method is employed in calibration. The standard reference samples are provided by National Technical Committee for Standardization of Microbeam Analysis (China) or produced by SPI Supplies* (America). The analytical precision was better than 5%.

**PETROGRAPHY**

**Clinopyroxenite**

Medium- to coarse-grained clinopyroxenite is commonly composed of > 60 mol.% of euhedral columnar (up to 2 mm in length) clinopyroxene, as well as various amounts of anhedral biotite and K-feldspar, and minor accessory minerals mainly including apatite, magnetite and analcite (Figure 3). The oriented euhedral cumulate clinopyroxene are commonly seen as imbedded crystal hosted in biotite, titanite and K-feldspar (Figures 3A,B) forming poikilitic texture. Besides, most of the cumulus clinopyroxene is compositionally zoned. Except occurring as oikocryst, K-feldspars also occur as fine-grained crystals coexisting with analcite in between clinopyroxene crystals (Figure 3C). Occasionally, biotite constitutes up to 50 mol.% of the rock and becomes euhedral with size up to 3 mm thus forming biotite clinopyroxenite and even glimmerite (Figure 3D). Euhedral apatite (0.1–0.5 mm in diameter), magnetite (0.1–1.0 mm) and occasionally ilmenite (~0.1 mm) are presented as minor phases, and in some biotite rich area apatite can reach 15 mol.% to form apatite ores. Additionally, at some contact boundaries of minerals, magnetite is more well-formed than apatite but less than clinopyroxene (Figure 3C). Moreover, the presence of veins of pure analcite with width of 1–2 mm are ubiquitous in the clinopyroxenite (Figure 3E). These veins are commonly composed of (1) analcite; (2) K-feldspar with exsolved lamellae of albite, which are approximately right to the vein; and (3) biotite, occasionally hosting anhedral clinopyroxene inclusions. The clinopyroxene adjacent to these veins is euhedral and usually rimmed by a darker greenish corona of clinopyroxene.

**K-Feldspar Clinopyroxenite**

K-feldspar clinopyroxenite is transitional in term of mineral assemblage and crystal mode between syenite and clinopyroxenite. These rocks are dark reddish in color and relatively coarse-grained in hand specimen scale, and mainly consists of euhedral K-feldspar and clinopyroxene, with anhedral biotite. The modal proportion of the three main minerals vary from place to place. The euhedral tabular K-feldspar usually vary from 1 to 7–8 cm in length and are preferentially orientated. The occurrence of clinopyroxene and biotite are constrained in the interstices between K-feldspar crystals. They usually exhibit poikilitic texture similar to the clinopyroxenites, whereas minor titanite (1–2 mm) occurs as individual grain (Figure 3F). Minor apatite (<1 mm) also exists as interstitial phase. In addition, some veins with 1–2 mm width are filled by analcite are commonly present in the K-feldspar clinopyroxenite as well.

**Pseudoleucite Clinopyroxenite**

Pseudoleucite clinopyroxenite is characterized by leucocratic pseudoleucite or cluster of pseudoleucite embedded in the melanocratic matrix (Figure 4A), which are predominantly composed of biotite, clinopyroxene, and subordinate garnet, apatite, magnetite and K-feldspar. Most of the pseudoleucite crystals are generally euhedral to subhedral, with the typical shape of leucite with ~1 cm in diameter (Figure 4B).

Similar to some of the pseudoleucite reported (Gittins et al., 1980; Comin-Chiaramonti et al., 2009), the pseudoleucite in Yaojiazhuang consists of dominantly fine-grained K-feldspar (~50 mol.%, 5–40 µm), nepheline (~30 mol.%, 5–20 µm) and muscovite (~15 mol.%, 5–20 µm) intergrowth at the core and nearly pure fine-grained K-feldspar at the rim (Figure 4). The fringes of the intergrowth are vermicular or flame-like or...
feather-like and usually perpendicular to the boundaries of the pseudoleucite grains. Within some pseudoleucite, coarsening of euhedral prismatic to granular nephelines up to \( \sim 200 \, \mu m \) is observed coexisting with K-feldspar, whereas many of these nephelines have been altered to kaolin and corundum (Figure 4F). Some anhedral (sub-round to round) and zoned K-feldspars (hundreds of \( \mu m \)) occasionally exhibit lineation and present either across or at the edge of the pseudoleucites (Figures 4E,F). Adjacent to the K-feldspar lineation, shapeless intergrowth of nepheline with muscovite or plus analcite pools and patches usually occur. Within some pseudoleucite crystals, nephelines are totally replaced by analcites, even in the vermicular intergrowths, and in such places, the amount of muscovite also increased accordingly. Individual mineral inclusions are prevalent in the pseudoleucite, predominantly lamellar biotite (5 mol.%; tens of \( \mu m \)) and accessory celestine (\( \sim 10 \, \mu m \)), titanite (up to \( \sim 100 \, \mu m \)),apatite (tens of \( 10 \, \mu m \)), calcite (up to \( 100 \, \mu m \); Figure 5A) and magnetite (\( \sim 10 \, \mu m \)). Minor zoned clinopyroxene inclusions (several to \( > 100 \, \mu m \)) only occur as clusters and are rather rare (Figure 5B).

The matrix comprises mainly euhedral biotite (0.1–1 mm), clinopyroxene (0.1–1.5 mm) and garnet (0.01–1 mm). These phases contribute \( \sim 85 \, \text{mol.\%} \) of the total amount of matrix but the individual proportions for different phases vary from place to place. In most cases, columnar clinopyroxene and laminar biotite are showing a preferred orientation consistent with the boundary line of the globules. Clinopyroxene commonly shows complex zonation (Figure 5C), for example, a core of relics or clastic of clinopyroxene with visible zonation, rimmed also by zoned clinopyroxene (Figure 5D). Euhedral apatite (20–200 \( \mu m \)), subhedral to anhedral magnetite (<50 \( \mu m \)) and anhedral K-feldspar (10–300 \( \mu m \)) account for 15 mol.% of the matrix. Apatite crystals are usually enclosed by garnets (Figure 5E). Other minor phases include analcite (<30 \( \mu m \)), titanite (20–200 \( \mu m \)), muscovite (several micrometers), calcite (several

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**FIGURE 3** | Microphotos of clinopyroxenite. (A) Poikilitic texture with zoned clinopyroxene chadacrysts and biotite and K-feldspar oikocrysts (plane-polarized light). (B) Titanite oikocrysts (Back-scattered electron, BSE, image). There are also micro co-existing K-feldspar and analcrite at the upright part of the picture. (C) Zonation of K-feldspar oikocryst, and there are micro co-existing K-feldspar and analcrite at the upright part of the picture (BSE image). (D) Glimmerite (scanned photo of thin section). (E) A vein filled by analcrite (scanned photo of thin section). (F) K-feldspar clinopyroxenite showing poikilitic texture in the interstices between K-feldspar grains (scanned photo of thin section). Ap, apatite; Anl, analcrite; Bt, biotite; Cpx, Clinopyroxene; Kfs, K-feldspar; Mt, Magnetite; Ttn, titanite.
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FIGURE 4 | Photos of pseudoleucite clinopyroxenite. (A) Field photo. (B) A polished hand specimen. (C) A representative pseudoleucite and the mineral phases in both of the pseudoleucite and the matrix (BSE image). (D) Vermicular intergrowth of K-feldspar, nepheline and muscovite (BSE image). (E) Compositionally zoned K-feldspars clusters in a lineation (BSE image). (F) Coarse nepheline pseudomorphs in the compositionally zoned K-feldspars (BSE image). Cc, Calcite; Cls, Celestine; Grt, garnet; Ms, Muscovite; Ne, Nepheline; others as before.

micrometers) and ilvaite (∼10 µm). Micro intergrowth of nepheline or analcite with K-feldspar is also observed as shapeless interstitial phases. Euhedral granular analcites are found in clinopyroxenes with expansion cracks in the host minerals (Figure 5F).

Syenite
The syenite is pale red in color and consists of mainly oriented euhedral to subhedral K-feldspar (>70 mol.%, 0.1–3 mm), clinopyroxene (10–15 mol.%, 0.1–2 mm). Pegmatitic syenite is observed at the center of the syenite (Figure 2E), where the length of K-feldspar can be up to 10 mm. The K-feldspar crystals commonly host exsolved albite lamellae, and in those samples collected at the center of the syenite, zonation of K-feldspar is observed (Figure 6A). Anhedral clinopyroxene grains commonly show compositional zonation, and in some samples near the clinopyroxenite, they are in round shape with a diameter about 100 µm scattering among the K-feldspars (Figure 6B). In the syenite near the clinopyroxenite, small (<50 µm) acicular-shaped clinopyroxene crystals are observed in the hosted K-feldspar crystals (Figure 6C). Accessory phases include anhedral biotite (0.1–0.3 mm), analcite (tens of µm), apatite (∼100 µm), titanite (tens of µm), magnetite (several to tens of µm) and calcite (∼10 µm). The calcite usually co-exists with analcite and exsolved albite (Figure 6D). Anhedral amphibole surrounding clinopyroxene is occasionally observed in the samples from the center of the syenite (Figure 6E). Besides, analcite veins are usually observed in the syenite (Figure 6F). K-feldspar adjacent to the vein is euhedral and usually grows across the veins. Analcite lamellae is usually observed in these K-feldspar crystals.

Dykes
Mafic and syenitic dykes pervasively cut all the three major lithologic units in Yaojiazhuang complex. In the field, mafic dykes (commonly ∼10 cm wide) are greenish to dark gray in color. The mafic dykes are fine-grained, lithologically similar to the clinopyroxenite, consisting of mainly oriented euhedral clinopyroxene (<1 mm), biotite (<1 mm) and K-feldspar...
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FIGURE 5 | Photos of pseudoleucite clinopyroxenite. (A) Calcite inclusion (BSE image). (B) Anhedral zoned clinopyroxene clusters as inclusions in pseudoleucite (BSE image). (C) Complex zonation of clinopyroxene in the matrix (BSE image). (D) A core of relics or clastic of clinopyroxene with visible zonation, rimmed also by zoned clinopyroxene (BSE image). (E) Apatite enclosed in garnets (BSE image). (F) Euhedral analcites in clinopyroxenes with expansion cracks (BSE image). Abbreviations are as before.

(<1 mm) of almost equal quantities (~30 mol.%). Analcite is commonly present in the interstices to form networks (~1 mm in width) separating the samples into different lithological parts (Figure 7A). In these analcite-rich zones, more clinopyroxene and biotite is observed. Minor phases include apatite, magnetite, and titanite are present.

According to the spatial relationship, both early and late stage syenitic dykes (commonly around 10 cm wide) had been recognized (Figure 7B). Nevertheless, the two stage dykes are similar in mineral assemblage and defined as clinopyroxene syenite. K-feldspar is the main phase in both of these two kinds of dykes (>80 mol.%) and hosts exsolved albite thin lamellae. Subordinate minerals are composed of biotite and clinopyroxene. Anhedral biotite mainly occurs as interstitial phases between K-feldspar grains. Small needle-like clinopyroxene is widespread in the host K-feldspar and biotite, which is interpreted as a result of quenching. Accessory minerals includes titanite, analcite and apatite. In some samples, euhedral titanite can grow up to 1–2 mm (Figure 7C). The late stage dyke is composed of K-feldspar, clinopyroxene, biotite with considerable amount of albite. K-feldspar accounts for 70% modal and its size is generally around 1–2 mm in length except some crystals could be up to 2–3 cm (Figure 7D). Clinopyroxene (20 mol.%) can be divided into at least two generations, one is as euhedral to subhedral zoned phenocryst up to 0.6 mm (Figure 7E), the other is as needle-shaped inclusions in the matrix commonly less than 50 µm in length. Clinopyroxene pseudomorph filled by albite, muscovite and magnetite is also observed (Figure 7F). Subhedral to anhedral biotite (<2 mm) constitutes the rest 10% modal percent. Anhedral albite is observed co-existing with K-feldspar either as exsolved lamellae or in the interstices of clinopyroxene and biotite crystals (Figure 7E). Euhedral titanite and apatite are also present as minor phases.

RESULTS

Scanning Electron Microprobe

Elements mapping (Figure 8) shows that the pseudoleucites are mainly enriched in Si, Al, Na, and K but poorer in Fe and Ca
relative to the matrix, which should be caused by the appearance of different minerals. The concentrations of elements within the pseudoleucites from edges to cores are relatively homogeneous, except that the compositionally zonation of K-feldspars slightly lack in Si and K. The coarse nepheline pseudomorphs are more enriched in Ca, Na but deficient in Fe compared to other parts of the pseudoleucite.

Whole-Rock Geochemistry

The whole-rock major and trace element analyses of the studied syenite, clinopyroxenite, pseudoleucite clinopyroxenite, as well as syenitic and clinopyroxenitic dykes from Yaojiazhuang alkaline complex are given in Supplementary Table 1.

Syenites generally display higher SiO₂ (57.2–64.1 wt.%), Al₂O₃ (16.6–18.8 wt.%), Na₂O (1.3–4.9 wt.%) and K₂O (8.0–11.1 wt.%) contents but lower CaO (1.0–2.2 wt.%), MgO (0.1–0.7 wt.%), total Fe₂O₃ (1.9–5.5 wt.%), and P₂O₅ (0.03–0.2 wt.%). Their molar K₂O/Na₂O ratios vary largely, from 1 to 5.7. The chemical composition of the syenitic dykes resemble that of syenites, containing 58.9–61.2 wt.% SiO₂, 15.5–18.6 wt.% Al₂O₃, 1.0–5.4 wt.% Na₂O, 7.1–12.8 wt.% K₂O, and relatively lower CaO (0.3–2.5 wt.%), MgO (0.3–1.1 wt.%), total Fe₂O₃ (1.5–6.4 wt.%), and molar K₂O/Na₂O ratio (1.1–4.8). Different K-feldspar clinopyroxenite show relatively lower CaO (4–7.5 wt.%), MgO (1.4–4.8 wt.%), total Fe₂O₃ (6–9.4 wt.%), TiO₂ (0.6–1.4 wt.%), and molar K₂O/Na₂O ratio (1.1–1.3 wt.%).}

In general, clinopyroxenites and biotite clinopyroxenites are characterized by relatively higher CaO (15.2–18.15), MgO (9.6–12 wt.%), total Fe₂O₃ (11.4–21.2 wt.%), TiO₂ (1.6–1.7 wt.%), and P₂O₅ (1.9–3.4 wt.%) but lower SiO₂ (40.9–42.7 wt.%), Al₂O₃ (4–7 wt.%), Na₂O (0.5–0.9 wt.%), and molar K₂O/Na₂O (1.1–4.8). Differently, K-feldspar clinopyroxenite show relatively lower CaO (4–7 wt.%), MgO (1.4–4.8 wt.%), total Fe₂O₃ (6–9.4 wt.%), TiO₂ (0.6–1.4 wt.%), and molar K₂O/Na₂O ratio (1.1–1.3 wt.%). The clinopyroxenite dyke exhibit an moderate composition between (biotite) clinopyroxenite and K-feldspar clinopyroxenite (49 wt.% SiO₂, 11.6 wt.% total Fe₂O₃, 6.8 wt.% MgO, 7.8 wt.% CaO, 11.6 wt.% Al₂O₃, and 2 wt.% P₂O₅), indicating the cumulations of mafic minerals such as...
FIGURE 7 | Petrographic photos of dykes. (A) Clinopyroxenite dyke showing the analcite network (scanned photo of thin section). (B) Field photo showing the relationship between syenitic dykes and faults. (C) Coarse euhedral titanite in late-stage syenitic dyke (scanned photo of thin section). (D) Pegmatitic K-feldspar crystal in late-stage syenitic dyke (scanned photo of thin-section). (E) Euhedral zoned clinopyroxene with aegirine rim, also showing the albite exsolution in the matrix (BSE image). (F) Clinopyroxene replaced by muscovite, magnetite and albite (BSE image). Abbreviations are as before.

clinopyroxenite, apatite and magnetite in the formation of (biotite) clinopyroxenites, and progressively high proportions of felsic minerals like K-feldspar in K-feldspar clinopyroxenites. On the Harker diagram (Figure 9), the composition of (biotite) clinopyroxenites, K-feldspar clinopyroxenites, syenites and dykes exhibit significant linear correlations among many elements such as FeO, MgO and CaO, implying that the geochemical evolution of Yaojiazhuang samples from clinopyroxenites to syenites are mainly controlled by successive accumulation and subtraction of cumulate minerals.

The composition of pseudoleucite clinopyroxenite matrix resembles those of clinopyroxenites and is characterized by relatively high total Fe₂O₃ (21.1 wt.%) and CaO (15.9 wt.%), moderate MgO (7.1 wt.%), Al₂O₃ (7.3 wt.%), K₂O (4.4 wt.%), and P₂O₅ (1.6 wt.%) but low SiO₂ (38.9 wt.%) and Na₂O (0.5 wt.%). On the contrast, the pseudoleucite contains higher SiO₂ (57.0 wt.%), Al₂O₃ (23.8 wt.%), Na₂O (2.2 wt.%), and K₂O (11.5 wt.%) but lower CaO (0.8 wt.%), MgO (0.4 wt.%), total Fe₂O₃ (0.8 wt.%), and negligible P₂O₅. Both of the pseudoleucite and matrix have high molar K₂O/Na₂O ratios, which are 3.4 and 5.6 respectively. The pseudoleucite also displays a high LOI value (3.9 wt.%), which is dominantly contributed by the contents of H₂O⁺ (3 wt.%) and CO₂ (0.4 wt.%).

The trace element and rare earth element (REE) compositions of the Yaojiazhuang intrusion are given in Supplementary Table 1 and illustrated in chondrite-normalized and primitive mantle-normalized diagrams in Figure 10. In general, (biotite) clinopyroxenites have higher REE contents than syenites, and the matrix of pseudoleucite clinopyroxenite has the highest. K-feldspar clinopyroxenites contain moderate REE contents between (biotite) clinopyroxenites and syenites. For most REEs and trace elements, the contents in pseudoleucite are very low, which may be inherited from the extremely low partition coefficients in leucite. All the rock types are characterized by significant enrichment in highly incompatible elements and light REE. Notably on the spider diagram, (biotite) clinopyroxenites and K-feldspar clinopyroxenites show a variation of P from positive anomaly to negative anomaly, symmetrical about that of the clinopyroxenite dyke, implying the fractionation/cumulation of apatite to form apatite ore hosted in clinopyroxenites.
The (biotite) clinopyroxenite and the matrix of pseudoleucite clinopyroxenite exhibit a distinct enrichment in high field-strength element compared to syenite and pseudoleucite. The pseudoleucite is significantly enriched in large ion lithophile elements (LILE), such as K, Rb, Ba, and Sr, commonly higher than those of syenites. In the REE patterns, all of rock types have similar slopes, whilst the concentrations of REE in clinopyroxenites and the pseudoleucite clinopyroxenite matrix are much higher than other samples. Despite two syenitic dykes, other samples do not show distinct Eu anomaly, with $(\text{Eu/Eu}^*)_N$ values generally vary around 1.

**Mineral Chemistry**

Silicates

Silicate minerals in Yaojiazhuang complex include clinopyroxene, K-feldspar, biotite, muscovite, garnet, nepheline, analcite, amphibole, apatite and titanite, and their compositions by EMPA are given in Supplementary Table 2.

Clinopyroxenes in all lithologies of the Yaojiazhuang intrusion generally show zonings with a Mg-rich core and an Fe-rich rim (Figure 11A). Zoned clinopyroxene has a composition of $\text{Wo}_{34.1-47.4}\text{En}_{34.7-42.8}\text{Fs}_{9.3-15.7}\text{Ac}_{1.0-4.2}$ in the core and to $\text{Wo}_{33.3-44.8}\text{En}_{20.5-29.5}\text{Fs}_{20.5-28.8}\text{Ac}_{5.2-15.3}$ in the rim (Figure 11A). Rims are usually become rich in Na, indicated by the high Ac endmember. Interestingly, the clinopyroxenes in both of pseudoleucite and the matrix of pseudoleucite clinopyroxenite are similar in composition as well. A compositional profile of a clinopyroxene crystal with oscillatory zonings in clinopyroxenite is shown in Figure 12A. Aegirine is widely recognized in the late-stage clinopyroxene syenite dykes with a composition of $\text{Wo}_{11.1}\text{En}_{6.1}\text{Fs}_{40.5}\text{Ac}_{40.2}$. A profile of a zoned clinopyroxene in this syenite dyke is shown in Figure 12C with an augite core and aegirine rim.

K-feldspars in the syenites in the center of the complex and the late-stage syenite dykes exhibit a relatively large compositional rang of $\text{Or}_{52.6-74.0}\text{Ab}_{24.9-46.5}\text{An}_{9.3-1.0}$. In contrast, K-feldspars in other parts of the Yaojiazhuang complex
are almost pure orthoclase, showing restricted compositional variation of Or$_{86-2-95}$Ab$_{4.8-13-8}$An$_{0.0-0.3}$ (Figure 11B). For example, the compositions of K-feldspar in both of pseudoleucite and matrix of pseudoleucite clinopyroxenite are almost pure orthoclase. Notably, albite is observed coexisting with K-feldspar as exsolved lamellae in clinopyroxenite, syenite and syenitic dykes.
FIGURE 10 | Primitive mantle normalized trace element spider diagram of (A) syenite, (biotite) clinopyroxenite and K-feldspar clinopyroxenite; (C) syenitic and clinopyroxenitic dykes; (E) pseudoleucite and matrix of the pseudoleucite clinopyroxenite; chondrite normalized rare earth element (REE) pattern of (B) syenite, (biotite) clinopyroxenite and K-feldspar clinopyroxenite; (D) syenitic and clinopyroxene dykes; (F) pseudoleucite and matrix of the pseudoleucite clinopyroxenite.

Published data of Yaojiazhuang complex are from Mu and Yan (1992) and Chen et al. (2013). The primitive mantle and C1 chondrite normalizing values are from Sun and McDonough (1989).

with composition Or$_{0.4-1.4}$Ab$_{91.6-99.3}$An$_{0.3-7.2}$. The most An-rich feldspar (An = 7) is found in the syenites in the center of the complex, where that zonation of K-feldspar with an Or-rich core and an Ab-rich rim is commonly observed (Figure 12B).

The composition of biotite in the Yaojiazhuang complex show restricted range, with FeO$_{tot}$ from 18.2 to 23.6 wt.%, MgO from 10.4 to 14.2 wt.%, TiO$_2$ from 1.8 to 2.5 wt.%, and K$_2$O from 9.4 to 9.8 wt.%. Accordingly, the biotite can be classified into the siderophylite group on the Fe/(Fe + Mg)–ΣAl diagram (Figure 11C).

All the analyses of garnets are Fe- (24.9–25.4 wt.%) and Ti-rich (1.6–4.8 wt.%) and occupy a limited compositional range...
FIGURE 11 | Compositional variation of major minerals from Yaojiazhuang complex. (A) Clinopyroxene; (B) K-feldspar with exsolved albite lamellae; (C) biotite; (D) garnet; (E) TiO$_2$–FeO–Fe$_2$O$_3$ ternary system (after Buddington and Lindsley, 1964). Data of clinopyroxenes and garnets from Fanshan ultrapotassic complex (Hou et al., 2015) are plotted for comparison.

on the Alm + Sp–Gro–Py diagram (Coleman et al., 1965; Figure 11D), which are essentially grossular (61.2–61.7%) and almandine (37.3–37.6%). However, compared to those from Fanshan complex unit 3 clinopyroxenite, whose FeO content commonly below 22 wt.% and TiO$_2$ above 5 wt.% (up to 12.3 wt.%), these garnets are richer in FeO but poorer in TiO$_2$.

Amphiboles found in both of clinopyroxenite and syenite are classified as ferro-pargasite, which is characterized by the significant high content of FeO$_{tot}$ (20.1–20.1 wt.%). The contents of SiO$_2$, TiO$_2$, MgO, CaO, Na$_2$O, and K$_2$O are 39.6–40.1 wt.%, 1.1–1.5 wt.%, 8.5–9.5 wt.%, 10.4–10.5 wt.%, 2.8–3.2 wt.%, and 1.9–2.0 wt.%, respectively. Notably, the amphiboles in our samples also contain 0.8–0.9 wt.% F.

Apatites are nearly pure fluorapatite characterized by $X_F$ [= molecular F/(F + Cl)] very close to 1.00 with high F (2.1–3.4 wt.%) relative to low Cl contents (<0.02 wt.%), which implies that they may crystallized from a high temperature, i.e., magmatic environment (Tacker and Stormer, 1989). Other oxides contents in all three lithology units exhibit no obvious differences, with...
SiO$_2$, CaO, and P$_2$O$_5$ contents are 0.4–1.0 wt.%, 50.6–53.1 wt.%, and 38.5–41.4 wt.% respectively.

The composition of nepheline is relatively identical in all pseudoleucite samples, the SiO$_2$, Al$_2$O$_3$, Na$_2$O, and K$_2$O contents of which are 41.5–42.6 wt.%, 34.2–34.5 wt.%, 15.8–16.1 wt.%, and 5.9–7.5 wt.%, respectively, while its CaO content is extremely low, which is 0.1–0.6 wt.%. Analcites are characterized by containing high sodium and water contents, with Na$_2$O contents are 11.3–12.5 wt.%, and total values range from 89.4 to 92.4 wt.%. The SiO$_2$ (52.2–56.3 wt.%), Al$_2$O$_3$ (22.8–25.7 wt.%) and K$_2$O (0.2–0.5 wt.%) contents vary limitedly.

Muscovites from all of the lithologies do not show significant difference, and show SiO$_2$ contents from 43.8 to 45.3 wt.%, Al$_2$O$_3$ from 34.9 to 38.2 wt.%, K$_2$O from 9.0 to 11.2 wt.%, FeO from 0.4 to 3.9 wt.%, MgO up to 1.6 wt.% and total values generally between 93 and 96 wt.%. The SiO$_2$, TiO$_2$ and CaO contents of titanite are 28.9–30.0 wt.%, 34.4–38.0 wt.%, and 25.8–27.4 wt.%, respectively, together with 0.2–0.4 wt.% F and a few Cl.

### Discussion

**Oxides**

Oxides in Yaojiazhuang complex mainly include magnetite and ilmenite, and the analyses of them are reported in Supplementary Table 3.

Except the FeO content of one magnetite from the matrix of pseudoleucite clinopyroxenite is relatively low [81 wt.% with Fe$^{3+}$(Fe$^{3+}$ + Fe$^{2+}$) = 0.57], others from both of syenite and clinopyroxenite all exceed 90 wt.% [90.3–93.8 wt.% with Fe$^{3+}$(Fe$^{3+}$ + Fe$^{2+}$) = 0.65–0.66]. The TiO$_2$ content of magnetite varies largely, from 0.3 to 5.2 wt.%, whereas no systematic trend can be found among them. SiO$_2$ content lies in 0.04–1.3 wt.%. In the TiO$_2$–FeO–Fe$_2$O$_3$ ternary system (Figure 11E; Buddington and Lindsley, 1964), magnetite from all lithologies are defined as titanomagnetite.

Ilmenite displays higher TiO$_2$ content (62.2 wt.%) and lower FeO content (28.0 wt.%). In the TiO$_2$–FeO–Fe$_2$O$_3$ ternary system (Figure 11E; Buddington and Lindsley, 1964), the ilmenite is yielded as ferropseudobrookite.

**Sulfates**

Sulfate, i.e., celestine is observed only in the pseudoleucite, and its composition is listed in Supplementary Table 4. Celestine contains mainly sulfur and strontium, with SO$_4$ and SrO contents are 39.76 and 49.92 wt.%. Both of SiO$_2$ and Al$_2$O$_3$ contents are 0.3 wt.%. Celestine contains nearly no F and Cl.

**Assessment on Alteration**

Most syenitic rocks have LOI values less than 1 wt.%, except one syenite (YJZ08-1) and a pegmatitic syenite dyke (YJZ04-1) which are 2.5 and 1.5 wt.%, respectively. The high LOI values for the two samples are mainly attributed to the bound water H$_2$O$^+$ and their moisture water contents show no significant differences to other samples (H$_2$O$^+$ = 0.1–0.4), thus we suggest that the high LOI values are due to a relatively high content of hydrous minerals like analcite and biotite. Similarly, the biotite clinopyroxenite YIJZ07-5, clinopyroxenite dyke (YIJZ07-2) and K-feldspar clinopyroxenite (YIJZ04-4) also show high contents of LOI values up to more than 3 wt.%, whereas their high LOI values are mainly constituted by high bound water up to 2.7 wt.%, consistent with the high contents of biotite and analcite on petrography. The pseudoleucite also has high LOI, high H$_2$O$^+$ (3 wt.%) but relatively low H$_2$O$^-$ (0.5 wt.%), owing to the widespread existing of muscovite, analcite and biotite. Hydrothermal alteration or weathering usually introduce water mobile elements, such as LILE, into rocks, thus the high contents of these elements may indicate the alteration process. However, either on the Sr vs. LOI (Figure 13A) or moisture water H$_2$O$^-$ diagram (Figure 13B), no significant correlation is observed for both of clinopyroxenitic or syenitic rocks, implying their primary geochemical signatures are preserved.

There are also classic indices to evaluate the degree of chemical weathering of silicate rocks. For instance, Plagioclase Index of Alteration (PIA) established by Fedo et al. (1995) is widely employed in evaluating the alteration of plagioclase- and/or...
K-feldspar-rich rocks, and Bases to Alumina ratio (BasesAl; Coleman, 1982) is a power tool for mafic rocks. The higher the PIA or the lower the BasesAl, the higher the degree of alteration. There is still no obvious correlation is present for synetetic rocks on the Sr vs. PIA diagram (Figure 13C) or for clinopyroxenites and K-feldspar clinopyroxenites on the Sr vs. BasesAl diagram (Figure 13D). The K-felspar clinopyroxenite with low BasesAl values show higher Sr contents than clinopyroxenites, which should be resulted from the higher proportions on Sr-bearing minerals like K-feldspar instead of alteration. Thus, their high Sr contents are able to represent their primary high LILE contents.

Crustal Contamination

Silica saturation index (SSI) was introduced by Motoki et al. (2010) to verify the role of crustal contamination by syenitic magma. The presence of pseudoleucite, nepheline and analcite indicate that the parental magma of Yaojiazhuang complex ought to be peralkaline and undersaturated in silica. The country rocks of the Yaojiazhuang complex is the Archean TTG, which are basically SiO₂-oversaturated and meta- to peraluminous in composition (e.g., Xie et al., 2019; Chen et al., 2020). Since the Yaojiazhuang complex were emplaced into these TTG, it seems unavoidable for the parental magmas were contaminated. Hence, the elevation of SSI is expected due to assimilation of TTG. Meanwhile, such interaction between parental magmas and wall rocks could also decrease the alkali content in the magmas as low-alkali affinity of the TTG (Motoki et al., 2015). Accordingly, on the SSI vs. molar (Na + K)/Al diagram (Figure 14A), there are four out of six syenites plotted on the left-top quadrant, implying most of syenites had been largely affected by crustal contamination. This inference is further supported by the plot of SSI vs. K₂O/(K₂O + Na₂O) diagram (Figure 14B).

Composition of Parental Magma

As stated above, the clinopyroxenite, K-feldspar clinopyroxenite and syenite exhibit coherent evolution trends both mineralogically and geochemically, indicating these lithologies have a common parental magma. Such inference is also supported by the similar Sr-Nd isotopic compositions of rocks from different lithologies (Chen et al., 2013). Whereas the intrusive contacts between these lithofacies suggest that they are not derived from a single batch of magma.
Therefore, it is more likely that there should be a deeper magma chamber where fractional crystallization of parental magma takes place.

Conventionally, the parental magma of Yaojiazhuang complex is thought to be ultrapotassic (Hou, 1990b; Mu and Yan, 1992; Chen et al., 2013), which is defined as $K_2O > 3$ wt.%, $MgO > 3$...
wt.%, and molar K$_2$O/Na$_2$O > 2 in whole-rock geochemistry according to Foley et al. (1987). While, the MgO contents for all of our syenite samples are below 3 wt.%, and those MgO enriched clinopyroxenite samples contain K$_2$O less than 3 wt.%. Despite that the molar K$_2$O/Na$_2$O in some samples can reach more than 5, there are still some syenite and clinopyroxenite samples having smaller K$_2$O/Na$_2$O ratios.

However, the definition in Foley et al. (1987) is based on the whole-rock compositions of sequences of volcanic rocks which are deemed to be liquids. However, rocks from Yaojiazhuang complex are cumulates, the whole-rock composition of which are controlled by the modal proportions of cumulate minerals. Thus, the low MgO K-feldspar-dominated syenites are expected to have low MgO content, and the low K$_2$O but MgO enriched clinopyroxene dominated clinopyroxenites are expected to be lower in K$_2$O but higher in MgO. Nevertheless, a feeder dyke (sample YJZ07-2) which is fine-grained and both ultrapotassic in composition (MgO = 6.8 wt.%, K$_2$O = 7.4 wt.%, molar K$_2$O/Na$_2$O = 3.5) could represent the composition of parental magma. Such inference is consistent with abundant presence of K-rich phases including biotite and K-feldspar in the complex. This dyke represents dominant phase among the basic dykes in the studied area and exhibits similar mineralogy and coherent variation trend geochemically with other dominant phases, i.e., clinopyroxenite and syenite. The moderate geochemical characteristics of the clinopyroxenite dyke between clinopyroxenites and K-feldspar clinopyroxenites indicate that clinopyroxenite and K-feldspar clinopyroxenite are derived from the cumulation of mafic minerals, and the syenite is a more evolved phase differentiated from them.

**Fractional Crystallization**

The bimodal clinopyroxenitic and syenitic series has been speculated to be formed via silicate liquid immiscibility (Hou, 1990b). The magma immiscibility mechanism has also been proposed to explain the petrogenesis of similar bimodal complex like in Hamilton Montana (Lelek, 1979). However, evidence from mineralogy is not consistent with the immiscibility model. For example, equilibrated immiscible Fe- and Si-liquids ought to crystallize same mineral assemblage but with different proportions (e.g., Vekslar et al., 2007). According to this criterion, this is not the case for the clinopyroxenite and syenite in Yaojiazhuang as albite is absent in clinopyroxenite and garnet is exclusively seen in the matrix of pseudoleucite clinopyroxenite. Major phase in fractionation crystallization or cumulation can be confirmed by geochemical data plotting. For instance, the Zr/Sr vs. Sm/Yb (Figure 14C), Dy/Yb vs. SiO$_2$ diagram (Figure 14D) and Sr vs. Ba diagram (Figure 14E) all suggest that a clinopyroxene dominant fractional crystallization/cumulation. Besides, the coupled decrease in Ba and Sr also suggest that the fractionation of K-feldspar plays a dominant role in the differentiation of syenitic rocks, which had also been illustrated by the positively correlation between K$_2$O/(K$_2$O + Na$_2$O) and SSp (Figure 14B). Combining the rock and mineral textures as well as geochemical characteristics, the crystallization sequence is inferred as in **Figure 15**, clinopyroxene/apatite/oxides → biotite/titanite → leucite/K-feldspar/analcite/aegirine → albite.

K-feldspars crystallized as primary phases from the melt including the euhedral crystals in syenite and those interstitial phases in between clinopyroxenes in clinopyroxenite. In addition, some K-feldspar could also be formed by break down of leucite in pseudoleucites. Analcite can be precipitate either as primary mineral in the intergrowths with K-feldspar or secondary phases replacing leucite and/or nepheline.

**Fluid-Mineral Interaction**

**Formation of Pseudoleucite**

Pseudoleucite is defined as a result of disintegration of Na-rich leucite into an intergrowth of K-feldspar and nepheline with addition of other mineral phases like muscovite, analcite etc., and it usually occurs as large megacryst in alkaline complexes globally (e.g., Viladkar, 2010). However, its genesis is enigmatic and remains a hotly debated topic (e.g., Knight, 1906; Bowen and Ellesstad, 1937; Larsen and Buie, 1938; Barrer and Hinds, 1953; Fudali, 1963; Seki and Kennedy, 1964; Hamilton and McKenzie, 1965; Davidson, 1970; Taylor and MacKenzie, 1975; Gittins et al., 1980). Several genetic models had been proposed including: (1) breakdown of early-formed leucite into K-feldspar and nepheline under subsolidus conditions (T = ~600°C) during late magmatic stage. (2) reaction of primary leucite with a Na-rich liquid. (3) introduction of Na$^+$ from hydrous glass or aqueous vapor by ion exchange. Here, we will briefly discuss these theories as the possible mechanism to produce the pseudoleucites in Yaojiazhuang complex.

Conventionally, problem arises to the subsolidus breakdown of leucite because although experiments shows that leucite could accommodate about 40 wt.% NaAlSi$_2$O$_6$ in the solid solution (Taylor and MacKenzie, 1975), natural leucite has never been observed to contain excess amount of Na to form the intergrowths with sodium-bearing phases on decomposition.
The abundant sodium-rich phase like nepheline and analcite in the mineralogy of the pseudoleucite suggest they are generated from a pre-existing Na-rich phase. Therefore, the pseudoleucite in Yaojiazhuang samples cannot be explained by subsolidus break down of potassic leucite alone, but processes introducing sodium into leucite to form a metastable sodic leucite prior to breakdown seem to be necessary (Taylor and MacKenzie, 1975).

Reaction between leucite and Na-rich melt/liquid was proposed as mechanism to produce a nepheline-K-feldspar pseudomorph and former leucite disappears (Bowen and Ellestad, 1937; Edgar, 1984). An outstanding feature of this theory is that, the leucite crystal morphology can hardly be well preserved during the reaction with melt/liquid (Taylor and MacKenzie, 1975). We observed in our samples, the pseudoleucite generally exhibit the pentagonal icositetrahedron morphology of leucite in macro scale, whereas many of their rims are curved and modified in micro scale, suggesting that such a reaction process cannot be totally ignored (Figure 4).

Indeed, the pseudoleucite is considered as a late-stage phase after mafic minerals due to these vermicular nepheline-feldspar or analcite-feldspar intergrowths are also present as anhedral interstices in the matrix or clinopyroxenites. The Na content in zoned clinopyroxenes are also observed progressively increase from core to rim, supporting to an increasingly sodic crystal mush in the late stage. Moreover, various minor mineral phases (e.g., biotite, clinopyroxene, apatite, titanite) occur within the pseudoleucite and show no obvious compositional differences to those in the matrix, suggesting they are appeared to be primary magmatic inclusions. These phases introduce additional elements into the pseudoleucites, which advises investigations only based in traditional petrogeny's residua system (SiO\textsubscript{2} into the pseudoleucites, which advises investigations only suggesting they are appeared to be primary magmatic evolution are the transformation of initial leucite to analcite in the matrix (Figure 5F), the widespread albite lamellae in K-feldspars and the pegmatitic syenites at the top of the intrusion (Figure 2E). In summary, the presence of pseudoleucite and hydrous primary minerals provided solid evidences for the activities of fluids. Fluids saturation and exsolution may have been achieved during decompression of replenished magmas from deep and during crystallization of anhydrous minerals like clinopyroxene.

**Implications for the Formation of the Apatite and IOA Ores**

The widespread interstitial biotite in clinopyroxenite and presence of glimmerite imply that the late-stage melt could be significantly hydrous. Dissolved aqueous fluid could be oversaturated at some point and exsolved from the crystallizing magma during decompression (e.g., Moore et al., 1998; Dixon et al., 1995; Newman and Lowenstern, 2002; Ballhaus et al., 2015) or during isobaric crystallization of anhydrous minerals (Candela, 1997), such as clinopyroxene and leucite in this study. Moreover, the presence of magmatic calcite also indicates that the parental magma could also be CO\textsubscript{2}-rich (Mu et al., 1999), which suppresses the H\textsubscript{2}O solubility in melt, resulting in exsolution of H\textsubscript{2}O at a relatively lower concentration (Caricchi et al., 2018; Edmonds and Woods, 2018). Hence, in the case of Yaojiazhuang, the scenario involving exsolution of aqueous fluids during the magmatic stage is expected. Besides, these aqueous fluids should also be abundant in F, due to F-rich biotite is widespread throughout the complex.

Experimental studies indicate that phosphate saturation in silicate melts is mainly controlled by SiO\textsubscript{2} and CaO, and addition of H\textsubscript{2}O and/or F in silicate magmas leads to saturation of apatite (Tollari et al., 2006, 2008). The Yaojiazhuang complex is characterized by relatively high CaO and low SiO\textsubscript{2} content. Hence, it is thus expected that the magmatic liquids during the differentiation of the complex may reach high concentrations in both Fe and P. Once the concentration of H\textsubscript{2}O and/or F was elevated by fractionation or other processes, saturation of apatite will be reached. Magnetite will also crystallize induced by decrease in P concentration in the melt (Tollari et al., 2006), rather than elevation of oxygen fugacity during the late stage of differentiation. In other words, apatite and magnetite preferentially co-crystallized from the magmas. Besides, the exsolution of aqueous fluids is considered as a potential mechanism in the formation of porphyritic or spheroidal structure rocks hosting economic ore deposits (Huber et al., 2012; Ballhaus et al., 2015; Guo and Audétat, 2017). As described above,
the IOA ores occurred as rocks with spheroidal structure in the field. Therefore, we propose that the fluids exsolution could also be a viable mechanism to concentrate not only magnetite but also apatite by wetting (Ballhaus et al., 2015) and formed the apatite and IOA ore in Yaojiazhuang complex. Notably, Knipping et al. (2015a; 2015b) have already advocated that flotation, coalescence and transport of magnetite-bubble pairs within a magma chamber is the key process for the formation of IOA ores. However, their model has only focused on enrichment of iron oxide instead of iron oxide plus apatite, i.e., IOA ores. Nevertheless, we admit that indeed, the role of exsolved fluids could be more important than we thought. Hence, if this is the case, the occurrence of apatite and IOA ores in Yaojiazhuang provides direct petrologic evidence for such a model involving fluids exsolution and concentration of iron oxide and apatite.

CONCLUSION

(1) The bimodal lithologies are more likely to be formed via crystallization fractionation of an ultrapotassic parental magma instead of silicate liquid immiscibility. Fractionations of both clinopyroxene and K-feldspar, and repeated magma replenishment in the crustal chamber played an important role during the differentiation of primitive ultrapotassic magmas. Crustal contamination may also have been involved during the solidification of the complex. Except that, during the emplacement of Yaojiazhuang magma, decompression, fractionation of anhydrous clinopyroxene and the enhancement by CO2 may result in the exsolution of an aqueous fluid.

(2) The genesis of pseudoleucite is related to original leucite crystallization from an evolved, silica-undersaturated magma followed by the interaction with sodium-rich fluids during the late-magmatic stage. Transportation of aqueous fluid within crystal mush is believed to be a key factor in generating the leucite pseudomorph with ambiguous crystal boundary.

(3) The differentiation under fluid-rich condition in the crust could be responsible for the petrogenesis of Yaojiazhuang ultrapotassic complex, thus the concentration of apatite and IOA ores is presumed to be closely related to the fluid exsolution process as well.

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DATA AVAILABILITY STATEMENT

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

AUTHOR CONTRIBUTIONS

TH and RP designed the research, conducted the field trips, and all the analysis works together. Following data treatment were undertaken by RP and TH. All authors actively participated in the subsequent discussions and interpretation of the data as well as in the preparation of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/feart.2020.00357/full#supplementary-material
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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