Current theories suggest that the first continental crust on Earth, and possibly on other terrestrial planets, may have been produced early in their history by direct melting of hydrated peridotite. However, the conditions, mechanisms and necessary ingredients for this crustal formation remain elusive. To fill this gap, we conducted time-series experiments to investigate the reaction of serpentinite with variable proportions (from 0 to 87 wt%) of basaltic melt at temperatures of 1,250–1,300°C and pressures of 0.2–1.0 GPa (corresponding to lithostatic depths of ∼5–30 km). The experiments at 0.2 GPa reveal the formation of forsterite-rich olivine (Fo90–94) and chromite coexisting with silica-rich liquids (57–71 wt% SiO2). These melts share geochemical similarities with tonalite-trondhjemite-granodiorite rocks (TTG) identified in modern terrestrial oceanic mantle settings. By contrast, liquids formed at pressures of 1.0 GPa are poorer in silica (∼50 wt% SiO2). Our results suggest a new mechanism for the formation of the embryonic continental crust via aqueous fluid-assisted partial melting of peridotite at relatively low pressures (∼0.2 GPa). We hypothesize that such a mechanism of felsic crust formation may have been widespread on the early Earth and, possibly on Mars as well, before the onset of modern plate tectonics and just after solidification of the first ultramafic magma ocean and alteration of this primitive protocrust by seawater at depths of less than 10 km.

Keywords: experiment, basaltic melt-serpentinite rock interaction, TTG, ophiolites, Hadean eon, Noachian, Mars, protocrust
INTRODUCTION

The conditions and mechanisms that led to the production of the earliest intermediate to felsic (Si- and Al-enriched) crust on Earth and Mars are the subject of intense debate (Rudnick and Gao, 2003; Harrison, 2009; Reimink et al., 2014, 2016; Sautter et al., 2015, 2016; Burnham and Berry, 2017). There are some indications that this crust was composed of igneous rocks of granitic composition (I-type) and may have been formed in a tonalite-trondhjemite-granodiorite (TTG)-like environment, i.e., by partial melting of a garnet-bearing lower crust, as far back as the geological record goes (e.g., the parental melt of the 4.37–4.02 Ga Jack Hills detrital zircon crystals, Burnham and Berry, 2017). Ancient granodiorite rocks have also been identified on Mars (Sautter et al., 2015; Sautter et al., 2016), indicating that formation of felsic crust occurred on other terrestrial planets even without plate tectonics and subduction. In this context, melting of dry peridotite at shallow depths does not produce appropriate intermediate to felsic melt compositions comparable to that of bulk continental crust (Hirschmann et al., 1998; Rudnick and Gao, 2003; Harrison, 2009; Reimink et al., 2014, 2016; Burnham and Berry, 2017). As a result, most currently accepted models for the generation of felsic crust on the Earth or Mars consider a complex multi-stage process at different depths, which involves extraction of basaltic magma from peridotitic mantle followed by fractional crystallization (Reimink et al., 2014, 2016; Udry et al., 2018) and/or re-melting of hydrated mafic rocks at lithospheric conditions of >0.6 GPa (Harrison, 2009; O’Neil and Carlson, 2017). However, specific conditions during the Hadean lead us to envisage petrological mechanisms that are not typical of present-day felsic magma generation (i.e., not involving subduction; see Herzberg et al., 2010; Sautter et al., 2016). For example, the early terrestrial crust of intermediate to felsic composition might have been created by direct melting of a serpentinitized (e.g., hydrated) peridotite at shallow depths (Rudnick, 1995; Rudnick and Gao, 2003). A possible present-day analogue is the formation of tonalites-trondhjemites in the shallow mantle beneath oceanic spreading centers (called oceanic plagiogranites in this context) (Coleman and Peterman, 1975; Amri et al., 1996; Amri et al., 2007). Tonalite-tonalite veins are common features throughout the cores in the few sites of the Deep Sea Drilling Project, and Ocean Drilling, Integrated Ocean Drilling and International Ocean Discovery Programs (DSDP, ODP, and IODP) where the deep oceanic crust has been drilled along present-day spreading ridges. These rocks have been ascribed to local remelting processes in the presence of hydrothermal fluids along high-temperature normal shear zones. These rocks are not abundant (no more than 1% of the volume of the sections), but their abundance might be underestimated due to poor core recovery in the fault zones (e.g., Pietranik et al., 2017). Their origin is enigmatic, although understanding the origin of TTG rocks situated in modern oceanic lithosphere may open new perspectives on the formation of the first planetary felsic crust.

Rapid low-pressure melting experiments (≤1.0 GPa) involving peridotite and low water contents (up to 0.5 wt% in the bulk source) indicate that peridotite-derived partial melts may become enriched in SiO₂ (up to 57 wt%) and other lithophile elements (Al, alkalis), with abundances approaching those of continental crustal rocks (Hirschmann et al., 1998; Ulmer, 2001). The increase in the silica contents of partial melts at low pressures is generally considered to be due to the combined effects of water, alkalis, and low pressure on the structure of the aluminosilicate melt, shifting the olivine-pyroxene cotectics to higher SiO₂ contents through a decrease in the silica activity coefficient (Kushiro, 1968; Hirschmann et al., 1998; Lundstrom, 2000).

As an alternative, or complement to direct hydrous melting of peridotite, a currently untreated question is to what extent the presence of basaltic melt may have an impact on the production of felsic crust. This impact may be direct (through chemical reactions with the host peridotite) or indirect (by providing a heat source). In this respect, experiments that reproduce basaltic melt-peridotite reaction at low pressures (<0.8 GPa) are currently limited to systems undersaturated in aqueous fluid (e.g., Fisk, 1986; Kelemen et al., 1990; Morgan, and Liang, 2003; Van den Bleeken et al., 2010; 2011). Although the experiments show the presence of dry basaltic to andesitic-basaltic melts in association with olivine, no direct data exist concerning the partial melting of hydrated (serpentinized) peridotite (or serpentinite) triggered by the emplacement of basaltic melts at shallow crustal depths (5–15 km). This latter scenario is of primary interest in early Earth history (and possibly also for Mars), because the ultramafic-mafic protocrust formed from the crystallization of a magma ocean was most likely hydrated, either by volatiles released by the magma ocean itself or through interactions with liquid water available on the planetary surface (Albarède and Blichert-Toft, 2007; Elkins-Tanton, 2012). A present-day analogue can be observed at the level of the mantle/petrologic Moho boundary in the oceanic lithosphere (at depths of 6 km), where interactions of basaltic magmas with peridotites produce chromitite-dunite associations (Kelemen et al., 1995; Arai, 1997; Borisova et al., 2012; Zagtrdenov et al., 2018; Rospabé et al., 2019) and serpentinized peridotite (hydrated residual peridotite) undergoes partial melting (Benoit et al., 1999). Furthermore, such shallow conditions of serpentinite melting in the presence of basaltic melt might also occur in mantle plumes (Bindeman, 2008; Reimink et al., 2014, 2016; Borisova et al., 2020a), in Hadean heat-pipe volcanoes (Moore and Webb, 2013) or Hadean protorift volcanoes (Capitanio et al., 2020) and/or during meteorite impacts (Marchi et al., 2014).

The above considerations provide a strong motivation for the present study, which aims to directly investigate the generation of intermediate to felsic crust on young terrestrial planets. For this purpose, we have conducted a set of time-series experiments in serpentinite-basalt systems under relatively low pressures (≤1 GPa), corresponding to depths of ≤30 km. Modern (i.e., post-Archean) magmatism near the petrologic Moho in the oceanic lithosphere can then be used as a ground-truth test within its limits.
FIGURE 1 | Upper panel: Scheme of time-series runs at 0.2 GPa, showing starting configurations and final products containing felsic glasses. Three types of experiment are described: serpentinite dehydration, and mixed and hybrid runs. Ol, high-Mg olivine; Opx, orthopyroxene; Cpx, clinopyroxene; Chr, chromite or chromiferous magnetite; felsic glass, intermediate to felsic glasses and fluid, aqueous fluid. Lower panel: Backscattered electron images of the hybrid and mixed runs at 0.2–1.0 GPa pressures. (A) mixed run SB7 at 0.2 GPa; (B) zoom on the SB7 with magnification of 1,100; hybrid runs: (C) P32 at 0.2 GPa, (D) P1 at 0.5 GPa, (E) P3 at 1.0 GPa. Mixed samples (SB7) is homogeneous sample with high-Mg olivine and interstitial felsic melt (L intoxic). Hybrid samples (P32, P1, and P3) contain two zones: olivine-rich zone (the former serpentinite zone) with interstitial felsic melt (L intoxic) and zone of reacting basalt (L react). Ol, high-Mg olivine; Opx, orthopyroxene; Chr, chromite, and F, aqueous fluid. It should be noted that the homogeneous olivine-rich zone containing high-Mg olivine, chromite, interstitial felsic melt, and aqueous fluid is formed in the mixed run SB7 (R = 0.25) at an early stage (in less than 2 h), suggesting early attainment of steady state in this system with a high starting proportion of serpentinite (80 wt%). All details may be found in Table 1 and Supplementary Table S2.
EXPERIMENTAL STRATEGY AND METHODS

Experimental Strategy

To investigate the serpentinite rock-basaltic melt reactions, we have conceived three types of time-series experiments: (i) pure serpentinite dehydration in a basalt-free system, and so-called (ii) “mixed” and (iii) “hybrid” experiments in the serpentinite-basalt system as a function of pressure and basalt-to-serpentinite ratio. (Figures 1A–E; Table 1; Supplementary Tables S1, S2). The “mixed” experiments were performed on well-homogenized serpentinite and basaltic powders compressed together, whereas the “hybrid” experiments were performed using cylindrical pieces of serpentinite rock placed in the upper part of the capsule and a basaltic powder in the lower part of the capsule in contact with the base of the serpentinite cylinder (Figures 1A–E). These two distinct experimental setups were designed to simulate two principal geological scenarios of basaltic melt emplacement. The mixed experiments (with variable basalt-to-serpentinite mass ratios, $R_{bas/serp}$ or $R$ from 0.25 to 4) simulate the case of efficient fertilization of peridotite in order to achieve a closer approach to equilibrium and obtain stable associations of the minerals and melts produced by the reactions on the relatively short time scales that are accessible experimentally. The hybrid experiments (with generally higher $R$, from ~3 to ~7) were designed to simulate the process of basaltic melt infiltration into serpentinite rock, that does not necessarily reach equilibrium.

All experiments were conducted using two types of piston cylinder and an internally heated pressure vessel for durations ranging from 1 min to 120 h at pressures of 0.2, 0.5, and 1.0 GPa and temperatures of 1,250 and 1,300°C (see Supplementary Materials for details). In modern oceanic settings, temperatures of ~1,050°C are considered to be sufficient to initiate the reaction of serpentinized peridotite with basaltic magma at 0.2 GPa (Borisova et al., 2012). In the present experimental study, however, we chose to use higher temperatures (1,250–1,300°C) since these provide a closer analogue of the thermal conditions of the Hadean and early Martian upper mantle (Herzberg et al., 2010; Sautter et al., 2016). Redox conditions expressed relative to the conventional quartz-fayalite-magnetite buffer and serpentinite in the capsule. Note that our run durations (~48 h) are too short to allow use of a double-capsule technique with common mineral buffers of oxygen fugacity that require longer times to attain equilibrium (Matjuschkin et al., 2015). The redox conditions in our experiments were buffered by the initial $Fe^{2+}/Fe^{3+}$ ratios imposed by the starting basaltic glass and serpentinite in the capsule. Note that our run durations (≤48 h) are too short to allow use of a double-capsule technique with common mineral buffers of oxygen fugacity that require longer times to attain equilibrium (Matjuschkin et al., 2015). The redox conditions (i.e., oxygen fugacity, $f_O2$) during the runs are estimated from the compositions of co-existing olivine and chromite using equations of Ballhaus et al. (1991) that yield $ΔQFM$ values of +1.8 to +4.6 (where $ΔQFM$ denotes the log$_{10}$ value relative to the quartz-fayalite-magnetite buffer, Supplementary Table S2). Additionally, the $Fe^{2+}/Fe^{3+}$ ratios in several glass run products were derived from XANES (X-ray absorption near edge structure) spectroscopy using the Fe K-edge at the European Synchrotron Radiation Facility (ESRF) in Grenoble, France (Supplementary Figure S1). The $Fe^{2+}/Fe^{3+}$ ratios obtained by XANES correspond to $ΔQFM$ (+1.8 to +3.6) based on model of Borisov et al. (2018) (Supplementary Table S2), in excellent agreement with the values estimated from phase composition. The relatively oxidized conditions inferred for our runs are principally the result of the presence of water, although partial $H_2$ loss through the capsule wall cannot be fully excluded. However, redox potential does not have a significant influence on the concentrations of the major redox-insensitive elements of the felsic crust (predominantly Si, and Al).

We used the law of mass conservation and the minimization code of Oliphant (2006) (version 1.17). Two steps were applied to calculate phase proportions obtained in hybrid runs. In the first step, the initial MORB glass composition was introduced for the compositions of $L_{bas}$ (basaltic liquid) and $L_{int}$ (interstitial felsic liquid) (Supplementary Tables S1, S2). In the second step, the
basaltic and interstitial glasses (L_{bas} and L_{int}, respectively) were distinguished as different phases. The run products (glasses and minerals) were analyzed for chemical compositions (Supplementary Tables S3, S4; Supplementary Material) to make reasonable comparison with natural felsic melts and magmas. Furthermore, thermodynamic modelling was performed to constrain the equilibrium compositions of the liquids, minerals as well as fluid produced during serpentine dehydration and responsible for the felsic melt production (Supplementary Tables S5–S7; Supplementary Material).

Analytical and Microanalytical Methods
The bulk-rock chemical compositions of the starting serpentine and basalt samples have been analyzed by inductively coupled plasma-mass spectrometry (ICP-MS) and inductively coupled plasma-optical emission spectroscopy (ICP-OES), using a method developed at the SARM (Nancy, France). Scanning electron microscope (SEM) and electron probe microanalysis and mapping (EPMA) of minerals and glasses were performed at the Géosciences Environnement Toulouse (GET, Toulouse, France) laboratory and at the Centre de Microcaractérisation Raimond Castaing (Toulouse, France). Secondary ion mass spectrometry (SIMS) was applied to measure the H$_2$O contents at CRPG-CNRS (Nancy, France). Laser ablation-inductively coupled plasma-mass spectrometry (LA-ICP-MS) was applied to analyze major and trace element contents at the Max Planck Institute for Chemistry, (Mainz, Germany). XANES spectroscopy was used to estimate iron redox state in selected experimental glass products from Fe K-edge XANES spectra acquired at the FAME beamline of the European Synchrotron Radiation Facility (ESRF, France). All details of the applied methods may be found in the Supplementary Material.

Modelling Approaches
To interpret the experimental and analytical data obtained in this study and to enable comparisons with natural systems, we employed complementary modelling approach. Note that there is no currently available thermodynamic model and associated software tools allowing to directly simulate the aqueous fluid-assisted melting of serpentine in the presence of basaltic melt at pressures below 1 GPa, that could predict, simultaneously and within the same thermodynamic framework, the compositions of fluid, melts and minerals. By contrast, robust thermodynamic models currently exist for i) hydrous melts and ii) aqueous fluid phase. Therefore, we conducted two types of thermodynamic modelling to better constrain the compositions of the different phases produced in the experiments: i) calculations of liquid and mineral phase proportions and compositions formed at different physical chemical conditions (temperature, pressure, T-P) using MELTS and Magma Chamber Simulator, MCS software and ii) calculations of the equilibrium aqueous fluid composition and major element speciation in the fluid phase to constrain the capacity of the aqueous phase to mobilize and fractionate major elements at the experimental T-P conditions from the initial materials (basalt and serpentine). All details of the modelling part are reported in Supplementary Material and Supplementary Tables S5–S7.

RESULTS
Experimental Sample Description
In this section, we describe the results of 23 experimental runs performed at 0.2–1.0 GPa and 1,250–1,300°C to investigate pure serpentine dehydration and serpentine-basalt interaction in the mixed and hybrid systems with different durations (1 min–120 h), different proportions of serpentine (13–100 wt%) and variable basalt-to-serpentine initial ratios (R = 0.25–6.5) in the starting material (Table 1; Supplementary Table S2).

Dehydration Experiments at 0.2–0.5 GPa
Three experiments on pure serpentine dehydration (100 wt% serpentine) at 0.2–0.5 GPa and 1,250°C were performed with different run durations. In experiment P29 (0.5 h duration), we observe a very porous aggregate consisting of enstatite [92–95 Mg# = 100 Mg/(Mg + Fe$^{2+}$)], high-Mg olivine (Fo$_{91-92}$, mol% forsterite) and chromite [69–73 Mg#, and 68–95 Cr# = 100 Cr/(Cr + Al), Figures 1A, 2, Supplementary Tables S2, S3]. Similarly, in the long-duration experiments P27 and P28 (48 h), the products are represented by a re-crystallized powder containing enstatite, high-Mg olivine and chromite. No trace of silicate glass was found in the run products. In all dehydration experiments at 0.2 and 0.5 GPa, fluid was present, likely produced by serpentine decomposition.

Mixed Experiments at 0.2 GPa
Based on textural and compositional characteristics, three types of mineral-glass assemblage can be distinguished in nine samples from the mixed experiments on serpentine-basaltic melt reactions with different proportions of serpentine (R = 0.25–4.0) in the starting material.

1) Most samples of the SB (Serpentine-Basalt) series with high initial proportions of serpentine (R = 0.25–1.0) are characterized by a homogeneous zone of polyhedral unzoned crystals of high-Mg olivine (Fo$_{90-94}$, mol%) associated with interstitial felsic glasses (SiO$_2$ = 62–71 wt%) as well as fluid bubbles (Figures 1A, B, 2). Chromite (Cr# = 68–89; Mg# 56–73) is associated with the euhedral olivine grains, and crystals of clinopyroxene (Mg# 58–68) are characteristic of the runs with R = 1. Figures 1A, B, 2 show that samples SB7, SBbis2, SBter3 with high initial proportions of serpentine (R = 0.25) are characterized by a single zone composed of polyhedral unzoned olivine (Fo$_{90-94}$) and chromite crystals associated with interstitial felsic glasses (L$_{inter}$) and interstitial crystals of chromite and fluid bubbles. Remarkably, the proportions of interstitial melt are high (33–40 wt%).

2) Experiments SB1, SB4, SBbis1, and SBter2, with lower initial proportions of serpentine (R = 1–4), are characterized by polyhedral unzoned olivine
TABLE 1 | Experimental run conditions and the resulting phases with the phase proportions.

| №  | Run   | P  | T   | Time | Percentage of the starting components (wt%) | R                          | Resulting experimental phases                        |
|----|-------|----|-----|------|---------------------------------------------|----------------------------|------------------------------------------------------|
|    |       | (GPa) | (°C) | (h)  | SERP | BAS* | BAS-to-SERP mass ratio |                                      | (Phase proportion in wt%) |
| 1  | P37   | 0.2 | 1250 | 0.5  | 100  | -    | 0                        | Ol (54.1) + Opx (30.4) + Chr (2.6) + F (12.8) |
| 2  | P28   | 0.2 | 1250 | 48   | 100  | -    | 0                        | Ol + Opx + Chr + F                          |
| 3  | P27   | 0.5 | 1250 | 48   | 100  | -    | 0                        | Ol (55.8) + Opx (28.6) + Chr (2.7) + F (12.8) |

Serpentinite dehydration experiments

| №  | Run   | P  | T   | Time | Percentage of the starting components (wt%) | R                          | Resulting experimental phases                        |
|----|-------|----|-----|------|---------------------------------------------|----------------------------|------------------------------------------------------|
| 1  | P33   | 0.2 | 1250 | 48   | 19.95 80.05 | 4.0 | Ol (39.6) + Cpx (22.3) + Chr (6.8) + L_int (34.0) + F (3.3) |
| 2  | P34   | 0.2 | 1250 | 48   | 94.89 5.11 | 1.0 | Ol + Cpx + Chr + L_int + F |
| 3  | P35   | 0.5 | 1250 | 48   | 19.95 80.05 | 0.25 | Ol (59.2) + Chr (6.5) + L_int (33.5) + F (5.9) |
| 4  | P36   | 0.5 | 1250 | 48   | 80.01 19.99 | 1.0 | Ol (37.9) + Cpx (19.3) + Chr (6.7) + L_int (39.9) + F (2.1) |
| 5  | P37   | 0.2 | 1250 | 120  | 28.2 71.8 | 1.0 | Ol (56.0) + Cpx (22.0) + Chr (7.1) + L_int (38.9) + F (2.1) |
| 6  | P38   | 0.2 | 1250 | 120  | 5     19.99 | 0.25 | Ol (57.5) + Chr (6.0) + L_int (35.0) + F (6.5) |

Mixed experiments

| №  | Run   | P  | T   | Time | Percentage of the starting components (wt%) | R                          | Resulting experimental phases                        |
|----|-------|----|-----|------|---------------------------------------------|----------------------------|------------------------------------------------------|
| 1  | P37   | 0.2 | 1250 | 0.5  | 28.2 | 71.8 | 2.5 | L bas (72.8) + L_int (0.4) + Ol (17.0) + Opx (6.8) + Cpx + Chr_Mgt (0.0) + F (3.0) |
| 2  | P38   | 0.2 | 1250 | 2.5  | 17.7 | 82.3 | 4.8 | L bas (72.7) + L_int (8.7) + Ol (10) + Opx (7) + Chr (0.0) + F (1.8) |
| 3  | P35   | 0.2 | 1250 | 5    | 18.8 | 81.2 | 4.3 | L bas + L_int + Ol + Opx + Chr + Chr_Mgt + Amph + F |
| 4  | P33   | 0.2 | 1250 | 48   | 19.8 | 80.2 | 4.1 | L bas (57.1) + L_int (15.4) + Ol (13.1) + Opx* (0.0) + Chr (6.0) + Amph (6.6) + F (1.7) |
| 5  | P42   | 0.2 | 1250 | 120  | 17.6 | 82.4 | 4.7 | L bas (~58) + L_int (~27.8) + Ol (12.8) + Chr_Mgt (0.0) + F (1.4) |
| 6  | P11   | 0.5 | 1300 | 0.02 | 19.2 | 80.8 | 4.2 | L bas + L_int + Ol + Opx + Chr + F |
| 7  | P15   | 0.5 | 1300 | 0.5  | 15.7 | 84.3 | 5.4 | L bas + L_int + Ol + Opx + Chr + Mgt |
| 8  | P18   | 0.5 | 1300 | 5    | 13.4 | 86.6 | 6.5 | L bas + L_int + Ol + Opx + Chr + Mgt + F |
| 9  | P36   | 0.5 | 1300 | 8    | 14.6 | 85.4 | 5.8 | L bas (~F) |
| 10 | P3  | 1.0 | 1300 | 2.5  | 15.3 | 84.7 | 5.5 | L bas + L_int + Ol + Opx + Chr_Mgt |
| 11 | P7   | 1.0 | 1300 | 9    | 15.9 | 84.1 | 5.3 | L bas |

*P, pressure; T, temperature; SERP, serpentinite; BAS, basalt; L_int, interstitial glass; L_bas, hydrous basaltic glass; Serp, serpentine minerals; Ol, olivine; Opx, orthopyroxene; Opx*, ferrosilite; Cpx, diopside; Amph, amphibole; Chr, chromite; Chr_Mgt, chromiferous magnetite.

Hybrid Experiments at 0.2 GPa

The samples from hybrid experiments with low starting proportions of serpentinite (18–28 wt%) have lower initial proportions of serpentinite (20 wt%) and a periphery (80–280 μm thick) consisting of olivine and interstitial felsic glass. In sample P32, which was kept for 2.5 h at 1,250°C and 0.2 GPa, the former serpentinite zone has a harzburgitic core (olivine Fo93±1 and orthopyroxene Mg# = 95 ± 3) and an outer thick rim composed of olivine and interstitial glass. Chromite is localized exclusively in the former serpentinite zone. By contrast, the intermediate-duration sample SBbis3 (5 h) is characterized by a residual olivine-rich zone, even though the starting material contains the same proportion of serpentinite as SBter1 (20 wt%).

3) Finally, the longest run sample SBter1 (48 h) with the lowest initial proportion of serpentinite (20 wt%) is represented by homogeneous basaltic glass formed by complete hybridization of the starting basaltic liquid with serpentinite. By contrast, the intermediate-duration sample SBbis3 (5 h) is characterized by a residual olivine-rich zone, even though the starting material contains the same proportion of serpentinite as SBter1 (20 wt%).

Hybrid Experiments at 0.2 GPa

The samples from hybrid experiments with low starting proportions of serpentinite (18–28 wt%; R = 2.5–4.7) produced at 0.2 GPa pressure show two distinct zones: a former serpentinite (or olivine-rich) zone and a zone of quenched basaltic melt (Figures 1C, 2; Table 1; Supplementary Tables S2, S3). The olivine-rich zone in sample P37 shows a core composed of enstatite (Mg# = 96.5 ± 2.1) and high-Mg olivine (Fo93±1), and a periphery (80–280 μm thick) consisting of olivine and interstitial felsic glass. In sample P32, which was produced after 2.5 h at 1,250°C and 0.2 GPa, the former serpentinite zone has a harzburgitic core (olivine Fo93±1 and orthopyroxene Mg# = 95 ± 3) and an outer thick rim composed of olivine and interstitial glass. Chromite is localized exclusively in the former serpentinite zone.
FIGURE 2 | Backscattered electron images and elemental X-ray maps of the hybrid and mixed runs to investigate aqueous fluid-assisted partial melting of peridotite at pressures of 0.2 and 0.5 GPa. The images show mineral, glass and fluid phase associations observed in the capsules, particularly the olivine-rich zones after quenching. Ol, Opx, Chr, Lint, Lbas, and F on the images stand for olivine, orthopyroxene, chromite, interstitial and basaltic glass, and fluid bubbles, respectively. The run numbers correspond to those used in Table 1; Supplementary Table S2. (P29) The serpentinite dehydration sample produced at 0.2 GPa displays a single zone represented by enstatite with high-Mg olivine and orthopyroxene. (SB7, SB bis2, and SB ter3) The mixed samples with highest initial proportions of serpentinite (80 wt%; \( R = 0.25 \)) are characterized by a unique polyhedral olivine-rich domain of unzoned crystals of olivine associated with interstitial felsic glass \( L_{\text{int}} \) and fluid bubbles. Chromite is associated with the euhedral olivine grains (high-Mg olivine \( F_{\text{O}2,\text{high}} \)). (P32) The hybrid run sample P32 lasting 2.5 h at 1,250°C and 0.2 GPa (\( R = 4.6 \)) shows a former-serpentinite zone with an inner harzburgite part and an outer olivine-rich part (olivine \( F_{\text{O}2,\text{low}} \)) composed of olivine and interstitial glass. Chromite is localized (Continued)
produced at 0.5 GPa and 1,300°C (except P36 at 1,250°C) with run durations from 1 min to 8 h (Table 1; Supplementary Tables S2, S4). In sample P1 with the shortest run duration (1 min), the quenched glass zone and former-serpentinite zone are both present. The former-serpentinite zone contains fine-grained aggregates (5–10 µm) of high-Mg olivine Fo95, enstatite with Mg# = 95, chromite (Cr# = 89) and an interstitial glass of basaltic andesite composition (Figures 1D, 2). Chromite crystals (a few microns in size) are disseminated within this zone. The bubbles in the quenched melt zone are evidence for the presence of fluid water during quenching. Sample P15 (0.5 h) also shows two distinct zones (quenched basaltic melt zone and olivine-rich zone). The quenched basaltic melt zone consists of hydrous basaltic glass characterized by a high magnesium content (11 wt% MgO). The former serpentinite zone consists of 5–20 µm aggregates of high-Mg olivine Fo93 and enstatite with Mg# = 97. These aggregates are associated with an interstitial glass of andesitic composition and chromiferous magnetite. The outer part of this zone is characterized by a layer (~120 µm thick), consisting of olivine crystals and interstitial glass. Chromiferous magnetite is concentrated in two large areas (up to 200 µm and 400 µm in size) or disseminated near these areas. Sample P18 (5 h of interaction) is also characterized by a hydrous basaltic zone and a former-serpentinite zone (Table 1; Supplementary Tables S2, S4). However, in this case, the olivine-rich zone consists of two areas; the outer area of nearly 200 µm width contains olivine and basaltic interstitial glass with disseminated chromite (a few microns in size), while the inner area is situated 200 µm from quenched basaltic glass. This inner area consists of high-Mg olivine Fo91, enstatite with Mg# = 97 and contains aggregate of magnetite (~20 × 40 µm). For the longest experiment of this series (P36, 8 h), serpentinite has completely dissolved into the basalt melt (Table 1; Supplementary Tables S2, S4). Run products show a homogeneous basaltic glass with high MgO content (11.5 wt%). More details are reported in Borisova et al. (2020a). Numerous bubbles with dendritic amphibole aggregates are present in the glass, suggesting amphibole growth upon quenching.

Hybrid Experiments at 1.0 GPa

Two experiments with low initial proportions of serpentinite (~15 wt%; R = 5.3–5.5) in the starting material were performed at 1.0 GPa and 1,300°C. The run products from experiment P3 (2.5 h) contain a basaltic glass zone and the former-serpentinite zone (Figure 1E; Table 1; Supplementary Tables S2, S4). High-Mg olivine (Fo92), enstatite with Mg# = 96 and chromiferous magnetite are associated with interstitial glass of basaltic composition (up to 51 wt% SiO2) in the former-serpentinite zone. A longer-duration experiment (P7, 9 h) yields products containing hydrous basaltic glass with 13.0 wt% MgO (Supplementary Tables S2, S4). More details are reported in Borisova et al. (2020a).

Summary of the Major Findings From the Whole Series of Experimental Data

1) Highly magnesian olivine (Fo91–92, mol% forsterite), orthopyroxene [92–95 Mg# = 100 Mg/(Mg + Fe²⁺)] and chromite [69–73 Mg#, and 68–95 Cr# = 100 Cr/(Cr + Al)] are found in the serpentinite dehydration runs containing only serpentinite as starting material (R = 0), but no interstitial glass can be detected (Figure 3A).

2) Bulk assimilation of the dehydrated serpentinite by the basaltic melt and production of homogeneous crystal-free hydrous basaltic melts (Lbas) are consistently observed in the hybrid and mixed experiments with the highest proportions of basalt and the lowest proportions of serpentinite (R ≥ 4, Borisova et al., 2020a) in the starting material and with durations longer than 5 h at pressures of 0.2–1.0 GPa (Figure 3B; Table 1; Supplementary Tables S2–S4).

3) The shortest hybrid runs at pressures of 0.5–1.0 GPa, and in all hybrid runs at 0.2 GPa lasting for 0.5–120 h contain olivine-rich zone composed of high-Mg olivine, Cr-rich spinels and interstitial glasses (Lint) (former-serpentinite zone or olivine-rich zone) and zone of hydrous basaltic glass (Figures 1C–E; Figure 2; Figure 3B; Table 1; Supplementary Tables S3, S4). The infiltration of basaltic melt into serpentinite at high basalt-to-serpentinite ratios (R = 2.5–6.5), as simulated in the hybrid experiments, is able to generate new interstitial melts of intermediate to felsic composition.
Silica contents ranging from 57 to 67 wt%) in most hybrid 0.2 GPa experiments. Only two shortest (≤0.5 h) of the 0.5 GPa runs yield interstitial melts of intermediate composition (SiO₂ = 56–60 wt%), whereas the shortest (2.5 h) 1.0 GPa run generates interstitial melt of mafic composition (SiO₂ = 50 wt%). These results are in line with the fact that the combined effects of water, alkalis and low pressure on the structure of the aluminosilicate melt shift the equilibrium olivine-pyroxene cotectics to higher SiO₂ contents (Kushiro, 1968; Hirschmann et al., 1998; Lundstrom, 2000).

4) Finally, a homogeneous and stable olivine-rich zone containing mostly high-Mg olivine (Fo 90–94, mol%), chromite (56–73 Mg#, and 68–89 Cr#) and interstitial felsic glass is systematically observed in all mixed (SB series) time-series experiments with low basalt-to-serpentinite ratios (R = 0.25–1.0) in the starting material, suggesting the formation of stable association of olivine, chromite and felsic melts at 0.2 GPa (Figures 1A,B, 2, 3A, and 4; Supplementary Table S3).

Interstitial Glass Composition and Equilibration With Minerals
In the hybrid and mixed runs at 0.2–1.0 GPa, the olivine-rich zones host interstitial pockets of glass ranging in size from 10 to 200 μm, showing enrichment in silica (up to 66–71 wt%), aluminum, alkalis and water (Figures 4, 5, Supplementary Tables S2–S4). The calculated chrome-olivine pair equilibration temperatures range from 1,222 to 1,261 °C (Supplementary Table S2), in accordance with the runs temperature of 1,250 °C, confirming an attainment of thermodynamic equilibrium between solid phases during experiments even if Fe-Mg partition coefficients between olivine and the associated felsic melt range from 0.19 to 0.24 in the hybrid runs and from 0.08 to 0.25 in the mixed runs according to model of (Toplis, 2005), indicating a lack of Fe-Mg equilibrium between phases involving the felsic liquid (although such low values as 0.16 have been reported in alkali-rich systems). This variability may be at least partly due to quench artifacts. Indeed, the glass transition temperature of hydrous andesitic
Fe-Mg disequilibrium. Hybrid run samples P32, P33, and P42 yield olivine-felsic glass Fe-Mg partition coefficients of 0.24 ± 0.13, close to the theoretical range of 0.31 ± 0.05 (Toplis, 2005) (Supplementary Table S2), suggesting that the felsic melts are close to Fe-Mg equilibrium with the associated olivine.

The major element composition of the glass produced in these experiments is mainly controlled by pressure, but also depends on the basalt-to-serpentinite initial ratio (R) in the mixed runs. The glasses produced in the hybrid run lasting 2.5 h at 1.0 GPa are mafic (50 wt% SiO₂), with an average MgO content of 10 wt%, similar to the glasses produced in previous studies of equilibrium partial melting of hydrated peridotite at similar pressures (Ulmer, 2001). By contrast, generally higher SiO₂ (57–71 wt%) and low basalt-to-serpentinite initial ratios (R = 0.25) produce dacite glasses with up to 70–71 wt% SiO₂ contents (Figures 4–6, Supplementary Table S3). Furthermore, the 0.2 GPa glasses are enriched in alkalis (up to 3.9 wt% Na₂O+ K₂O), aluminum (up to 20.4 wt% Al₂O₃) and chromium (up to ~710 Cr ppm) (Figures 5, 6; Supplementary Tables S3, S4).

The experimental glass composition obtained in the 0.2 GPa mixed experiments is in an agreement with that obtained through modeling using MELTS at condition of the melts saturation with H₂O fluid (Figures 5, 6; Supplementary Table S6). In the modeling, we used mixture of serpentinite with basalt in different proportions comparable to those observed in the mixed experiments (see Supplementary Material). The modeled phase proportions (e.g., melt proportions up to 35 wt %, Supplementary Table S6) and the mineral compositions are in agreement with those of the experimental ones (melt proportion up to 40 wt%, Supplementary Table S2). For example, high-Mg olivine (91–93 mol% Fo), chromite (20–45 mol% of chromite molecule), aqueous fluid and Si-rich liquid (up to 61 wt% SiO₂ on anhydrous basis) appear in equilibrium assemblages at 1,250–1,300°C, 0.2 GPa and ΔQFM+2 to +3 in the models using a mixture of 80–95 wt% serpentinite and 20–50 wt% basalt, respectively. The felsic liquids contain up to 67 wt% SiO₂ if the composition of aqueous fluid is taken into consideration (Supplementary Table S6). Additionally, the MELTS liquids modelled at 0.2 GPa (up to 61 wt% SiO₂ on anhydrous basis) are richer in SiO₂ than those modelled at 0.5–0.9 GPa (up to 55–59 wt% SiO₂ on anhydrous basis) (Supplementary Table S6), confirming our experimental observations at 0.5 and 1.0 GPa.

To evaluate the result of energy constrained interaction between serpentinite and basalt, we have performed Magma Chamber Simulator (MCS, Bohrson et al., 2014; Bohrson et al., 2020; Heinonen et al., 2020) modeling in the range from 0.1 to 0.5 GPa at condition of the magma saturation with H₂O fluid (Supplementary Table S7). The MCS modeled liquid composition is in excellent agreement with that of the experimental felsic melts (Figures 5, 6). The only exception is Mg index, but this is likely related to the Fe-Mg re-equilibration of...
the experimental liquids with the predominant olivine upon quenching.

**DISCUSSION**

**Felsic Melt Formation**

The measured glass compositions vary from basaltic andesite to dacite, which is consistent with the general tendency of olivine-saturated partial melts to become progressively richer in SiO$_2$ with decreasing pressure and increasing water content (e.g., Hirschmann et al., 1998). Additionally, it is widely known that water fluid expands the thermodynamic stability of forsterite-rich olivine (e.g., Green, 1973; Nicholls et al., 1973) and chromite (Veksler and Hou, 2020). Thus, it is not surprising that high-Mg olivine in association with chromite is recorded in equilibrium with felsic melts of dacitic composition in our low-pressure experiments (0.2 GPa). Most glasses produced at 0.2 GPa are richer in Si, Al, and alkalis than dry glasses produced at 0.1 MPa (Fisk, 1986) as well as nearly dry glasses produced at 0.5 GPa (Kelemen et al., 1995). The hydrous intermediate to felsic melts produced in our experiments are likely formed by the fertilization of olivine-rich zones due to the presence of aqueous fluids produced by the interaction of serpentinite and basaltic melt at 0.2–1.0 GPa. At lower pressures, this process is associated with the incongruent melting of harzburgite producing olivine at the expense of orthopyroxene. Indeed, according to Kushiro (1968), Shaw (1999), Shaw et al. (1998), Lundstrom (2000), Shaw and Dingwell (2008), and Borisova et al. (2020a) silica-rich liquids may be formed via orthopyroxene dissolution during melt-peridotite interactions. The presence of polyhedral and mainly unzoned olivine crystals, co-existing with homogeneous interstitial felsic melt in the olivine-rich zone of the longest runs (Figure 2; Table 1; Supplementary Table S2), supports the growth of new olivine that attains equilibrium with the melt at 1,250°C and 0.2 GPa.

A steady state is established within less than 2 h after the start of reaction in the mixed runs (Figures 3, 4). The fluid-present reaction leads to the formation of high-Mg olivine + chromite + felsic melt. Note that the composition of the interstitial felsic melt produced in this way is a function of the basalt-to-serpentinite mass ratio ($R$). High and constant SiO$_2$ contents (70–71 wt%,...
FIGURE 6 | Magnesium index (100 Mg/(Mg + Fe)), major and minor Ti, Al, P oxides (in wt%), and Cr (in ppm) vs. SiO₂ (in wt%) in interstitial glasses produced at 0.2–1.0 GPa during serpentinite-basalt interaction of hybrid and mixed runs. The major element compositions of the measured (meas.) melts are recalculated to 100% on a volatile-free basis. The error bars reflect the felsic glass inter-sample heterogeneity compared to the mean values (Supplementary Tables S3, S4). The glass composition is compared to the starting basalt (BAS) and serpentinite (SERP), the anhydrous 0.1 MPa experimental glasses (Fisk, 1986), modern TTG (plagiogranites) in mantle sections of ophiolites (Amri et al., 1996, 2007; Shervais, 2008; Xu et al., 2017), the bulk Earth continental crust (Bulk CC) (Wedepohl, 1995; (Continued)
observed in most hybrid and mixed experiments. The calculations and feldspar-like glass phases, are in good agreement with those orthopyroxene, together with minor amounts of clinopyroxene predicted to form at equilibrium are olivine and (olivine-rich zone, see above). The major mineral phases by serpentinite dehydration in the presence of basaltic melt is felsic glasses). Modeling shows that the aqueous analysis of the run products described above (e.g., interstitial qualitative support for our conclusions drawn from the exchange with olivine and chromite during quenching. (modelling of the element solubility in the aqueous lower the temperature of the serpentinite, the higher the SiO2 crystallization will be discussed in a future paper. iii) The increasing of all element concentrations in the fluid, in agreement with the increase of water solvent density and water solvation power (e.g., Pokrovski et al., 2013). A similar systematic order of decreasing concentration in the fluid is maintained over the whole range of pressures and basalt-to-serpentinite initial ratios: Si > Na > K > Al >> Fe > Ca > Mg. This finding strongly suggests that intrinsic uncertainties in the thermodynamic model are unlikely to obscure the large amplitude of general trends. The low concentrations of Fe and Ca (typically 100 to 10,000 times less abundant than Si, Na, K or Al), and in particular of Mg (10^6–10^8 less abundant than Si, Supplementary Table S5), are in agreement with the typical order of elemental abundance in natural hydrothermal-magmatic fluids of relatively low salinity as inferred from fluid inclusions (e.g., Kouzmanov and Pokrovski, 2012; references therein). This is corroborated by our experimental results, providing evidence for major transfer of Si, Na, K, and Al from basalt and serpentinite to felsic melts. Remarkably, the Si/Mg ratio (∼10^-6–10^-8) in the fluid phase predicted by calculations systematically increases by almost 1,000 times with decreasing pressure from 1 to 0.2 GPa (Supplementary Table S5). Even though our calculations are semi-quantitative at such extreme conditions for the fluid phase (see Supplementary Material), this large relative change is an additional indication that relatively low pressures should favor the enrichment of both fluid and melt phases in Si relative to “mafic” elements (Mg) in water-present systems, thus promoting the formation of felsic (i.e. Si-rich) melts quenched to glasses, as observed in our direct experiments.

Aqueous Fluid Composition

In all hybrid and mixed experiments at 0.2 and 0.5 GPa, fluid bubbles are present, likely produced by serpentinite decomposition. Therefore, we conducted thermodynamic modelling of the element solubility in the aqueous fluid phase (Supplementary Material), which provides an independent qualitative support for our conclusions drawn from the analysis of the run products described above (e.g., interstitial felsic glasses). Modeling shows that the aqueous fluid generated by serpentinite dehydration in the presence of basaltic melt is systematically enriched in Si, Al, Na, and K (Supplementary Table S5) compared to both the initial and reacted serpentinite (olivine-rich zone, see above). The major mineral phases predicted to form at equilibrium are olivine and orthopyroxene, together with minor amounts of clinopyroxene and feldspar-like glass phases, are in good agreement with those observed in most hybrid and mixed experiments. The calculations predict high-Mg olivine (Fo93–96 mol%) and orthopyroxene (Mg# 94–96) compositions, which are within the range analyzed in the hybrid run products (Supplementary Table S3, S4), suggesting local thermodynamic equilibrium was attained between the fluid and these minerals.

The aqueous fluid in mixed and hybrid experiments is predicted to be enriched in Si (up to 6 wt%), Na (up to 4 wt%), K (up to 0.4 wt%), and Al (up to 0.2 wt%), with other elements being much less abundant (Mg, Ca, Fe typically < ~10 ppm). Increasing the pressure from 0.2 to 1.0 GPa generally results in an increase of all element concentrations in the fluid, in agreement with the increase of water solvent density and water solvation power (e.g., Pokrovski et al., 2013). A similar systematic order of decreasing concentration in the fluid is maintained over the whole range of pressures and basalt-to-serpentinite initial ratios: Si > Na > K > Al >> Fe > Ca > Mg. This finding strongly suggests that intrinsic uncertainties in the thermodynamic model are unlikely to obscure the large amplitude of general trends. The low concentrations of Fe and Ca (typically 100 to 10,000 times less abundant than Si, Na, K or Al), and in particular of Mg (10^6–10^8 less abundant than Si, Supplementary Table S5), are in agreement with the typical order of elemental abundance in natural hydrothermal-magmatic fluids of relatively low salinity as inferred from fluid inclusions (e.g., Kouzmanov and Pokrovski, 2012; references therein). This is corroborated by our experimental results, providing evidence for major transfer of Si, Na, K, and Al from basalt and serpentinite to felsic melts. Remarkably, the Si/Mg ratio (∼10^-6–10^-8) in the fluid phase predicted by calculations systematically increases by almost 1,000 times with decreasing pressure from 1 to 0.2 GPa (Supplementary Table S5). Even though our calculations are semi-quantitative at such extreme conditions for the fluid phase (see Supplementary Material), this large relative change is an additional indication that relatively low pressures should favor the enrichment of both fluid and melt phases in Si relative to “mafic” elements (Mg) in water-present systems, thus promoting the formation of felsic (i.e. Si-rich) melts quenched to glasses, as observed in our direct experiments.

Compositional Features of Interstitial Glasses

The intermediate to felsic glasses from the hybrid and mixed runs were analyzed not only for major and minor (Cr, Ni) but also for trace elements (Supplementary Material). Figure 7 demonstrates incompatible trace element compositions (normalized to the primitive mantle) of the interstitial felsic glasses. The positive Pb anomaly relative to light rare earth
elements (LREE) observed in felsic glasses is an important signature of typical continental rocks compared to mantle-derived magmas (Rudnick and Gao, 2003). The incompatible trace element patterns of the felsic glasses are nearly identical to those of modern TTG (plagiogranite) rocks found in the mantle sections of ophiolites (Amri et al., 1996; 2007; Shervais, 2008; Xu et al., 2017). The felsic glasses produced in runs at 0.2 GPa have major and trace element contents (Si, Ti, Al, Mg, Ca, Na, K, P, and Cr, REE), as well as Mg/(Mg + Fe), that share some similarities with modern TTG situated in the mantle sections of ophiolites (Figures 5–7). The 0.2 GPa intermediate to felsic melts are characterized by lower SiO2 and Na2O + K2O contents and higher Cr contents compared with Archaean TTG rocks (Figures 5, 6). The slight enrichment in LREE of the felsic glasses produced in our experiments differs from the strongly fractionated patterns of Archaean tonalite-trondhjemite-granodiorite (TTG) suites (Moyen and Martin, 2012). This is in line with the conditions of shallow felsic crust formation by a mechanism distinct from the “sagduction” or plate subduction operating during the Archaean (Moyen and Martin, 2012). At similar SiO2 contents, the higher Mg index and Cr contents (up to 1,400 Cr ppm) in the experimental felsic glasses produced at 0.2 GPa contrast strongly with the 4.02 Ga Idiwhaa gneisses, northwest Canada (Figure 6), suggesting that the Hadean Acasta Gneiss Complex magmas did not form due to reaction of serpentinite with basaltic melts. Except for fluid-mobile B and Pb, the trace element compositions of the 0.2 GPa interstitial felsic glasses are similar to those of the starting basaltic melt (Figure 7). Additionally, hafnium contents (0.5–4.0 ppm Hf) and Lu/Hf ratios [LuN/HfN = 0.11–2.2, normalized to the primitive mantle composition of Sun and McDonough (1989)] in the 0.2 GPa felsic melts are similar to those of the starting basaltic melt (2.7 ppm and 0.7, respectively), suggesting that the experimental melts might have inherited their trace element and Hf isotope signatures from a precursor basaltic reservoir (Supplementary Table S3).

GEOLOGICAL SIGNIFICANCE

Shallow Felsic Crust Production by Melting of Serpentinitized Peridotite

Our study provides new insight into potential mechanisms of intermediate to felsic crust formation. Serpentinitized peridotites and mafic cumulates are the most abundant constituents of the modern slow-spreading oceanic lithosphere at the level of the Moho mantle-crust boundary (Cannat, 1993). Cyclic magmatic and hydrothermal processes in the modern Earth likely lead to the generation of intermediate to felsic material within the peridotitic lithosphere below spreading centers at high temperature and shallow (<10 km) depths. These processes include repeated intrusions of mantle-derived basaltic magmas (O’Hara and Mathews, 1981; Amri et al., 1996; 2007; Borisova et al., 2012), where reactions are possible between hot basaltic melts and a previously formed cold hydrated ultramafic lithosphere during periods of low magma supply and oceanic water circulation at shallow depth (Figure 8). The melting of serpentinitized peridotite produced in our experiments leads to the formation of dunite-wehlrite- or chromitite-type rocks (Figures 1–3A,B) similar to their natural analogues described in ophiolites (e.g., Dick, 1977; Benoit et al., 1999; Borisova et al., 2002c; Zagrtdenov et al., 2018; Rospabé et al., 2019; Borisova et al., 2020b). Our experimental data thus support an empirical model for the reaction between hydrated mantle and basaltic magma inferred from natural data on chromite-hosted inclusions from the Oman ophiolite (Borisova et al., 2012). The olivine-rich aggregates containing interstitial hydrous silica-saturated liquids are gravitationally unstable, like those obtained in our experiments. This is because of the high density contrast between hydrous felsic melts (<2.4 g/cm3, Supplementary Table S6) and mafic silicate minerals (3.2–3.3 g/cm3) and chromite (~5.0 g/cm3), while melt mobility is also favored by low viscosity. With reference to processes occurring at oceanic spreading centers (Amri et al., 1996, 2007; Benoît et al., 1999), which lead to the formation of
ophiolites, low-density hydrous felsic melts generated below and/or at the Moho transition zone are collected in decimeter to meter sized pods and veins which then progressively coalesce into larger intrusions in the shallow oceanic lithosphere. Alternatively, the felsic melts produced by aqueous fluid-assisted partial melting of peridotite may remain scattered in peridotites at shallow depths <10 km, as shown by olivine-hosted melt inclusions in peridotite xenoliths (Hirschmann et al., 1998). If segregation into intrusions takes place, cooling (e.g., due to hydrothermal circulation) will result in crystallization of these hydrous felsic liquids. In addition to the generation of low-temperature plagiogranite magmas in the oceanic crust due to partial melting of hydrated gabbros (Koepke et al., 2004), we suggest that high-temperature plagiogranites melts and magmas (or high-temperature TTG) may be formed through aqueous fluid-assisted partial melting of serpentinized peridotite.

Implications for the Early Earth
Hydrated peridotite, and its isochemical analogue serpentinite, has been suggested to be a major shallow component of the Hadean and Noachian protocrusts on Earth and Mars (Albarède and Blichert-Toft, 2007; Elkins-Tanton, 2012), being the result of the interaction of seawater-derived fluids with peridotites (Guillot and Hattori, 2013). Indeed, planets like Earth and Mars were massive enough to allow the formation of an early ultramafic protocrust during the earliest stages of planetary evolution. However, it remains unclear how early Hadean crustal rocks could have generated quartz-normative to quartz-saturated magmas of intermediate to
felsic composition. Direct natural evidence for the precursor of Hadean (≥4 Ga) felsic crust is scarce, and remains the subject of intense debate (Wedepohl, 1995; Rudnick and Gao, 2003; Harrison, 2009; Reimink et al., 2014, 2016; Burnham and Berry, 2017; Bell et al., 2018). Investigations of the 4.37–4.02 Ga Jack Hills detrital zircon crystals (JHZ), which are the oldest remnants of primordial felsic magmas on Earth, support the existence of a felsic crust during the earliest stages of the Hadean eon (Cavosie et al., 2006; Harrison, 2009; Burnham and Berry, 2017). Either reworking into younger crust or recycling of the Hadean crust into the mantle have been proposed to explain the lack of remnants of the earliest felsic crust on Earth (O’Neil and Carlson, 2017; Nédélec et al., 2017). However, studies of the Jack Hills zircon crystals allow to infer the presence of an igneous protolith that had experienced low-to-moderate temperature alteration by aqueous fluids, thereby ruling out a sedimentary source (Burnham and Berry, 2017; Whitehouse et al., 2017), although alternative opinions still exist (Harrison, 2009; Bell et al., 2018).

We propose that basaltic melts were introduced in the primitive ultramafic protocrust at the very end of the crystallization of the magma ocean, and/or by heat-pipe volcanoes through a stagnant lid (Moore and Webb, 2013), during the earliest stages of Earth evolution, or as proto-rifts at the end of the Hadean (Capitanio et al., 2020). Unfortunately, no traces of Hadean tectonic regimes survived so far. Thus, we propose here that the basaltic melts were intruded into an ultramafic protocrust, initiating dehydration processes. The released fluids promoted fertilization and partial melting of the peridotite, which in turn led to the production of tonalite-granodiorite magmas forming intrusive bodies in the peridotite protocrust at shallow depths <10 km (Figure 8). The segregation of felsic melts in the upper protocrust was favored by lithospheric hydraulic fracturing and the low density of the hydrous melts, with high melt proportions (up to 33–40 wt%, Supplementary Table S2) corresponding to high liquid volume fractions (φ) of ~37–45 vol%. Alternatively, the generation of such intermediate to felsic melts and magmas at shallow depths within the early ultramafic-mafic protocrust may also have been possible due to impact-induced melting (Marchi et al., 2014), particularly if the proportion of serpentinite was above 50 wt% in the hybrid system (R ≤ 1). Such melting would have favored intense mixing of shallow hydrated ultramafic and mafic components at high temperature (as modelled in our mixed experiments), well above the liquidus of basalt before the onset of modern plate tectonics (Figure 8).

Implications for Early Mars

The proposed mechanism of the aqueous fluid-assisted partial melting of peridotite induced by reaction with basaltic melt supports the possibility of felsic crust formation at depths of ≤10 km. The shallow depths of generation of felsic magma proposed in this study imply that the formation of silica-rich crust does not require convergent plate tectonics, in agreement with geodynamic simulations of the hotter mantle on early Earth and Mars (Herzberg et al., 2010; Sautter et al., 2016). The most favorable conditions for felsic crust formation are high-temperature interactions between serpentinized peridotite and basaltic melts at 0.2 GPa and low basalt-to-serpentinite mass ratio (≤1). Such conditions may have occurred in various geologic settings that do not require modern plate tectonics. Such conditions are highly probable at spreading centers, hot spots and Hadean heat-pipe volcanoes or proto-rifts. A felsic crust may also have been formed in a more transient way during intensive impact-induced melting of the ultramafic-mafic protocrust in the presence of a liquid water ocean on early Earth and Mars. Even though the geodynamic and thermal conditions on both planets have changed, small volumes of felsic crust are still forming by this process at modern spreading centers.

CONCLUSION

The proposed mechanism of aqueous fluid-assisted partial melting of peridotite induced by reaction with basaltic melt supports the formation of shallow felsic (continental) crust on other rocky planets such as Mars, provided liquid water or a volatile element-rich magma ocean was present at an early stage of their evolution. The finding of ancient serpentinites at the surface of Mars (Ehlmann et al., 2010) suggests that serpentinite formation and aqueous fluid-assisted melting of serpentinized peridotite may be envisaged during the earliest history of this planet. There is also evidence for a 4.43–4.13 Ga-old granodioritic and/or tonalitic crustal component on early Mars (Sautter et al., 2015; Sautter et al., 2016). These quartz-normative rocks, which are believed to represent the early Martian continental crust, cannot be produced from partial melting of the mantle and subsequent fractional crystallization (Sautter et al., 2015; Sautter et al., 2016). By contrast, recent modeling using the MELTS software (Udry et al., 2018) suggests that the felsic rocks of Gale crater were produced by the accumulation/fractionation of feldspar from basaltic melt; however, these authors make no comparison with modern TTG (plagiogranite) rocks. Our data demonstrated on Figures 5, 6 are also in line with our model and suggest that the Noachian granodiorite (felsic) rocks on Mars may have been produced by aqueous fluid-assisted partial melting of peridotite induced by reaction with basaltic melts. Survival of the primary intermediate to felsic liquids and formation of a solid crust may have been possible if the effects of gravitational overturn were limited (Elkins-Tanton, 2012; Bouvier et al., 2018). Thus, the primordial environment of the peridotite protocrust in the presence of mantle-derived (e.g., basaltic) melt at shallow depth is favorable for the production of silica-enriched melts provided liquid water is present in sufficient amounts. The mechanism suggested in our study may thus have led to formation of large volumes of the earliest felsic crust.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number can be found in the article/Supplementary Material, Supplementary Tables S1–S7.
AUTHOR CONTRIBUTIONS

ABo, AN, MT, and OS developed the conceptual idea of the study; ABo and NZ, SS prepared and conducted high T-P runs at BGI; NZ, OS, ABy, AT, VP, DV prepared and performed experiments at IEM; NZ, GP, WB, OM conducted thermodynamic modeling; GP, NZ, and AG carried out XAS and SIMS measurements, respectively; KJ, BS, and UW performed LA-ICP-MS analyses. IB performed bulk-rock analyses of serpentinite. ABo, NZ, SG, PP, and KC performed microanalytical measurements and mapping using EPMA; ABo, AN, GC developed geological applications; all authors contributed to data interpretation and manuscript writing.

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SUPPLEMENTARY MATERIAL

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

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