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Formation of the Flatreef deposit, northern Bushveld, by hydrodynamic and hydromagmatic processes

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Abstract

New lithological and whole rock compositional data show that the main platinum-group element (PGE) horizons of the Flatreef succession show strong compositional similarities to the Merensky and Bastard reefs of the western Bushveld Complex (WBC), notably in terms of many immobile and incompatible minor and trace elements such as TiO₂, Zr, Y, and REE. However, Al₂O₃, CaO, and Na₂O contents are markedly lower in the Flatreef, whereas MgO contents are higher. In view of broadly similar silicate mineral compositions in the Flatreef and the WBC reefs, we suggest that the major element compositional differences between the rocks are largely due to higher modal proportions of orthopyroxene and olivine and lower proportions of plagioclase in the Flatreef. The thickness of the mineralised interval is much greater in the Flatreef than in the WBC (several 10 s of m vs ~ 1 m) and the abundance of sulfides in the Flatreef is typically somewhat higher (on average ~ 1.5% vs ~ 1%). These data, complemented by textural observations, are interpreted to reflect enhanced hydrodynamic crystal sorting accompanied by percolation of sulfide melt through incompletely solidified cumulates. Further genetic constraints are provided by metal data: The concentration of Ni (~ 3000 ppm) in the Flatreef is broadly similar to that in the Merensky Reef of the WBC, but Cu contents are markedly higher (average ~ 1500 ppm vs 700 ppm in the WBC). The concentrations of most PGE are slightly lower (Flatreef ~ 1.5–2 ppm Pt, ~ 100–150 ppb Rh; WBC MR 3.7 ppm Pt, 240 ppb Rh), but Pd has broadly similar contents (Flatreef ~ 1.2–2 ppm; WBC MR 1.54 ppm). The relatively high Cu content of the Flatreef is interpreted as a result of assimilation of Cu sulfides from the sedimentary floor rocks. The reason for the enrichment of Pd relative to Pt, especially in the basal rocks, remains unclear. It could reflect mobilisation of Pd via a fluid phase from the country rocks or the interior of the intrusion, relatively enhanced partitioning of Pd into the sulfides, or relative Pt depletion of the earliest magma pulses in response to Pt alloy fractionation triggered by contamination with reducing country rocks.

Introduction

Grobler et al. (2019) used a ~ 700-km drill core database to document the stratigraphy of the “Flatreef”, a highly mineralised, flattened portion of the Platreef located down-dip of previous mining and exploration operations in the northern lobe of the Bushveld Complex (Fig. 1 in Maier et al. 2020). In contrast to the relatively poorly layered Platreef, the Flatreef shows more regularly layered reefs of the type found in the western and eastern Bushveld Complex (WBC and EBC), including the Merensky Reef (MR) and UG2 chromitite. These findings suggested that many of the processes responsible for the formation of the MR and UG2 equally controlled the formation of the Flatreef (Grobler et al. 2019). In the present paper, we use the extensive diamond drill core database from the Flatreef to place further constraints on these processes.
**Geological setting and igneous stratigraphy**

The Flatreef was intersected during exploration drilling on the farm Turfspruit, within the south-central sector of the northern limb, some 20 km to the N of Mokopane. The geological setting and igneous stratigraphy of the Flatreef have been presented in detail by Grobler et al. (2019) who correlated the most mineralised sequence of the Flatreef to the Upper Critical Zone (UCZ) of the WBC (Fig. 1). However, the Flatreef is much more mineralised (up to 4.5 ppm platinum-group element (PGE) over 90 m) than the Merensky and UG2 reefs of the WBC (each typically 5–8 ppm over ~1 m), and more contaminated resulting in abundant xenoliths and rafts of the floor rocks and reduced lateral continuity of layering.

The lowermost igneous rocks of the Bushveld Complex at Turfspruit consist of Lower Zone (LZ) ultramafic rocks that intruded, in a sill-like manner, into shales and dolomites of the Duitschland Formation. Next is the Flatreef, which is separated from the LZ by a screen of sedimentary rocks of variable thickness (metres to 100 s of metres). The basal Flatreef rocks typically consist of relatively poorly mineralised (<1 ppm PGE), strongly contaminated pyroxenite and norite. Thick intervals of calc-silicate may represent large xenoliths or floor remnants and roof pendants. This contaminated interval, termed the Footwall Assimilation Zone (FAZ), is overlain by the main mineralised interval which consists of a package of rocks containing interpreted correlatives of the UG2 chromitite and the Merensky and Bastard reefs. The recognition of this stratigraphic correlation at the mine matured over the course of several years of exploration. Thus, the ~1-m thick chromitite seam in the Flatreef was originally called “Main Chromitite” but has now been renamed UG2 chromitite (Grobler et al. 2019). The overlying main sulfide mineralised interval was initially called “T2” or “Main Reef” but is now referred to as Merensky Reef, and the uppermost mineralised interval was initially called “T1” or “Upper Reef”, but is now called Bastard Reef (Yudovskaya et al. 2017a; Grobler et al. 2019).

The Merensky Reef in the Flatreef succession typically comprises a basal Cr stringer, overlain by layers of pegmatoidal harzburgite termed M1L and/or pegmatoidal orthopyroxenite termed M1U (the subscripts L and U denoting “lower” and “upper”, but note that in some drill cores harzburgitic M1L can be above orthopyroxenitic M1U), a second Cr stringer, and medium-grained pyroxenite (termed M2), all of which may contain several percent PGE-rich sulfides (Fig. 2). The Merensky Reef is overlain by poorly mineralised or barren pyroxenite and/or norite (termed the Middling unit), and then by the Bastard Reef (Grobler et al. 2019).

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**Fig. 1** Proposed stratigraphic correlation of Flatreef with uppermost CZ in WBC (from Grobler et al. 2019)
Methods

In total, approximately 700 km of diamond drill core are available for study on the Turfspruit project. The deepest drill holes extend to >1800-m depth (Abernethy et al. 2020). Approximately 340 drill cores intersected the Flatreef, and 251,000 samples were assayed for Pt, Pd, and Rh as well as Au, Cu, Ni, Cr, and S. Lithophile major elements as well as a range of lithophile trace elements were determined for the reef intersection in 20 selected drill cores that are considered to represent a range of typical reef sequences (see Fig. 1 of Maier et al. (2020) for borehole locations). Rare earth elements were analysed in six of these drill cores (UMT 314, 336, 341D, 345, 378, and 390, 670 samples in total), and all six PGE (Os, Ir, Ru, Rh, Pt, Pd) were analysed in 116 samples. Average metal contents of BAR, M2, M1U, and M1L are listed in Table 1.

The concentrations of whole rock major elements were determined by XRF, S by LECO combustion, and base metals and precious metals by multi-acid digestion followed by ICP-MS and OES. Further details of the methods are outlined in Grobler et al. (2019). X-ray fluorescence (XRF) scanning microscopy was performed on the Maia Mapper laboratory XRF imaging system at CSIRO Mineral Resources, Clayton Victoria. The Maia Mapper (Ryan et al. 2018) comprises an Excillum D2 Liquid Metal Jet high brightness laboratory X-ray source (Larsson et al. 2011), a polycapillary X-ray focusing lens (www.xos.com), a 384-element Maia large solid angle (1.3 sr) X-ray detector array (Siddons et al. 2014) mounted in a coaxial back-scatter configuration, and a custom sample presentation and transport system designed for mineral samples up to 50 cm in length. The metal jet source was operated at 70 kV producing a broadband X-ray spectrum from its Ga-In-Sn liquid target, focused to an approximately 30-μm beam spot on sample. The sample was raster scanned in fly-scan mode with equivalent pixel sizes of 30×30 μm with full spectral data acquired at an effective 12 ms per image pixel. The quantitative multi-element images were produced using the dynamic analysis method with the CSIRO GeoPIXE software (Ryan 2000) maintaining the full spatial resolution for ~10 M pixel images for these samples. A standard sample manufactured by NIST (NIST 2012) was imaged between each mineral sample for calibration and fluence measurement and quality control. The procedures used to generate the false colour XFM element maps are described in Barnes et al. (2018).

Results

Petrography

Results of microXRF analysis of a coarse-grained Flatreef orthopyroxenite from the M2 reef in drill core TMT 006 are shown in Fig. 3. Approximate model proportions are up to ~80% orthopyroxene, ~10% clinopyroxene and plagioclase each, a few % sulfides, up to 1% phlogopite, and trace amounts of chromite. The image highlights a number of interesting textural features, including the mostly euhedral or subhedral habit of orthopyroxene, the subhedral habit of several of the large (oikocrystic) clinopyroxene crystals containing small orthopyroxene inclusions, and the location of the chromite crystals within interstitial plagioclase. The sulfides occur in patchy disseminated form in the interstitial space.
## Table 1: Average composition of PGE reefs in the Flatreef, compared with SARM bulk Merensky Reef sample from WBC

|          | BAR (n = 109) | M2 (n = 99) | M1u (n = 249) | M1L (n = 169) | MR SARM (n = 32) |
|----------|---------------|-------------|---------------|---------------|-----------------|
| SiO₂ %   | 53.39 ± 1.12  | 52.21 ± 3.98| 52.27 ± 1.96  | 45.29 ± 3.63  | 51.80           |
| Al₂O₃ %  | 5.70 ± 2.49   | 5.27 ± 2.04 | 5.10 ± 1.88   | 4.31 ± 1.48   | 8.30            |
| TiO₂ %   | 0.21 ± 0.03   | 0.23 ± 0.08 | 0.21 ± 0.08   | 0.14 ± 0.06   | 0.21            |
| MnO %    | 0.22 ± 0.03   | 0.22 ± 0.02 | 0.22 ± 0.03   | 0.21 ± 0.03   | 0.20            |
| CaO %    | 4.58 ± 1.28   | 4.11 ± 0.90 | 4.44 ± 1.92   | 3.49 ± 1.82   | 5.40            |
| MgO %    | 22.55 ± 2.87  | 23.43 ± 2.46| 23.86 ± 2.75  | 29.52 ± 3.65  | 20.10           |
| K₂O %    | 0.18 ± 0.10   | 0.19 ± 0.10 | 0.21 ± 0.14   | 0.17 ± 0.08   | 0.11            |
| Na₂O %   | 0.52 ± 0.25   | 0.39 ± 0.20 | 0.42 ± 0.25   | 0.26 ± 0.16   | 0.80            |
| Fe₂O₃ %  | 1.14 ± 0.20   | 1.25 ± 0.21 | 1.18 ± 0.19   | 1.54 ± 0.24   |                |
| FeO from Fe₂O₃ | 10.83 ± 1.52 | 11.53 ± 1.79| 11.24 ± 1.55  | 14.08 ± 1.94  | 10.15           |
| C %      | 0.13 ± 0.13   | 0.12 ± 0.08 | 0.12 ± 0.31   | 0.32 ± 0.29   |                |
| S ppm    | 7221 ± 3821   | 5642 ± 4412 | 8473 ± 4410   | 9135 ± 5539   | 4000            |
| LOI %    | 1.05 ± 1.21   | 1.14 ± 1.27 | 1.66 ± 1.53   | 5.25 ± 3.32   | 0.90            |
| Cr ppm   | 1676 ± 492    | 6859 ± 19,165| 4411 ± 8415   | 3538 ± 7043   | 6979            |
| Ni ppm   | 3029 ± 1349   | 2517 ± 1608 | 3357 ± 1438   | 3429 ± 1509   | 3100            |
| Cu ppm   | 1545 ± 851    | 1179 ± 835  | 1721 ± 769    | 1602 ± 926    | 700             |
| Pt ppm   | 1580 ± 1242   | 2218 ± 2770 | 1930 ± 1976   | 1253 ± 1096   | 3771            |
| Pd ppm   | 1254 ± 1035   | 1612 ± 1854 | 1959 ± 1702   | 1457 ± 1068   | 1543            |
| Au ppm   | 449 ± 391     | 272 ± 212   | 276 ± 191     | 194 ± 211     | 313             |
| % sulfide| 1.88 ± 1.09   | 1.41 ± 1.20 | 1.98 ± 1.25   | 1.97 ± 1.46   |                |
| Rh ppm (NiS) | 9 ± 82        | 166 ± 243   | 142 ± 157     | 128 ± 96      | 242             |
| Co ppm   | 118 ± 30      | 110 ± 35    | 138 ± 38      | 173 ± 43      | 143             |
| Ba ppm   | 46 ± 25       | 40 ± 20     | 55 ± 47       | 37 ± 15       | 50              |
| Sr ppm   | 62 ± 46       | 48 ± 31     | 50 ± 40       | 41 ± 18       | 75              |
| V ppm    | 124 ± 23      | 153 ± 93    | 151 ± 134     | 88 ± 24       | 60              |
| Rb ppm   | 9 ± 8         | 9 ± 5       | 12 ± 11       | 8 ± 4         | 9               |
| As ppm   | 24 ± 49       | 13 ± 6      | 15 ± 8        | 15 ± 7        | 2               |
| Y ppm    | 7 ± 2         | 7 ± 2       | 5 ± 2         | 5 ± 1         | 6²              |
| Zr ppm   | 13 ± 7        | 14 ± 8      | 15 ± 17       | 10 ± 9        | 15²             |
| La ppm   | 2.92 ± 1.51   | 2.73 ± 1.72 | 4.00 ± 4.44   | 3.20 ± 4.62   | 4.2             |
| Ce ppm   | 6.05 ± 2.89   | 5.54 ± 3.27 | 7.99 ± 8.60   | 6.08 ± 9.00   | 7.9             |
| Nd ppm   | 2.94 ± 1.32   | 2.94 ± 1.32 | 3.83 ± 3.67   | 2.94 ± 3.55   | 4.5             |
| Sm ppm   | 0.67 ± 0.34   | 0.75 ± 0.35 | 0.84 ± 0.62   | 0.74 ± 0.78   | 1.1             |
| Eu ppm   | 0.21 ± 0.06   | 0.24 ± 0.07 | 0.24 ± 0.09   | 0.22 ± 0.01   | 0.2             |
| Tb ppm   | 0.21 ± 0.05   | 0.21 ± 0.01 | 0.22 ± 0.06   | 0.24 ± 0.12   | 0.1             |
| Dy ppm   | 0.92 ± 0.38   | 0.90 ± 0.41 | 0.96 ± 0.48   | 0.78 ± 0.80   |                |
| Ho ppm   | 0.21 ± 0.06   | 0.24 ± 0.07 | 0.25 ± 0.17   | 0.24 ± 0.15   |                |
| Er ppm   | 0.67 ± 0.29   | 0.67 ± 0.22 | 0.69 ± 0.30   | 0.64 ± 0.48   |                |
| Tm ppm   | 0.20 ± 0.04   | 0.21 ± 0.01 | 0.21 ± 0.01   | 0.22 ± 0.04   |                |
| Lu ppm   | 0.20 ± 0.04   | 0.21 ± 0.01 | 0.21 ± 0.01   | 0.23 ± 0.08   | 0.1             |

**M1L** = harzburgitic pegmatoid in Flatreef, normally at base of the main reef, equivalent of pegmatoidal harzburgitic MR of Western Bushveld. **M1U** = orthopyroxenite pegmatoid in Flatreef, normally above M1L, equivalent of pegmatoidal orthopyroxenitic MR of Western Bushveld. **M2** = upper chroomite stringer plus overlying medium-grained pyroxenite of Flatreef, equivalent of non-pegmatoidal upper part of MR in Western Bushveld. **BAR** = upper reef in Flatreef, equivalent to Bastard reef in WBC. **SARM** = MR bulk sample in Western Bushveld

¹ Barnes and Maier (2002)
² Hutchinson et al. (2015)
showing early-formed sulfide-free oikocrysts, sulfides in face contact with the cumulus pyroxenes, and centimetre-scale domains where all the interstitial space is filled with sulfide separated by patches where none of it is.

**Average composition of the reefs**

In terms of lithophile elements, the PGE reefs of the Flatreef show certain compositional similarities to the Merensky Reef in the WBC, notably in terms of many immobile and incompatible minor and trace elements such as TiO$_2$, Zr, Y, and REE (Table 1, and Abernethy et al. 2020). However, the concentrations of the major elements Al$_2$O$_3$, CaO, and Na$_2$O contents are markedly lower in the Flatreef, whereas MgO and K$_2$O contents are higher than in the WBC Merensky Reef. The average sulfide content of the Flatreef is somewhat higher than in the Merensky Reef of the WBC (~ 1.5–2% vs ~ 1%). Nickel and Pd contents of the Flatreef are broadly similar to those in the MR of the WBC (~ 3000 ppm Ni, ~ 1.5–2 ppm Pd), but Cu contents are markedly higher (average ~ 1500 ppm vs 700 ppm) resulting in higher Cu/Ni and Cu/Pd, and Pt contents are lower (~ 2 ppm vs 3.7 ppm), resulting in lower Pt/Pd. Comparison of the 2 pegmatoidal units indicates that M1 tends to have mostly lower chalcophile (PGE, Cu) and incompatible elements (REE, Zr, Y, Rb, Ba, K$_2$O, Na$_2$O) than M1$_U$, but higher MgO, C, and LOI.

**Drill core profiles**

In the following section, we discuss in detail four drill cores that represent typical stratigraphic variants (UMT 263, 341d1, 378, and 449) (Fig. 4).

Drill core UMT 263 is located in Zone 1 (i.e., the relatively shallow northern portion of the property, see Fig. 1 of Maier et al. (2020)). The hanging wall sequence above the Bastard Reef (BAR) consists of interlayered pyroxenite, gabbro, and anorthosite. In this specific drill core, the hanging wall is unusually thick (~ 70 m) and contains relatively high PGE contents (frequently 10 to 100 s of ppb, occasionally up to 1 ppm), whereas most other hanging wall intervals of the Bastard Reef are PGE poor. Possibly, the interval represents a faulted block of rocks originally located below the Bastard Reef. This interpretation is consistent with the abundance of granite veins in this interval, likely along subsidiary fractures. The Bastard Reef consists of a ~ 5-m sulfide-bearing pyroxenite, overlying interlayered gabbro and anorthosite of the Middling unit with a sharp contact. M2 is represented by a thin (0.6 m) sulfidic pyroxenite. This is underlain by pegmatoidal harzburgite of M1$_L$, pegmatoidal orthopyroxenite (M1$_U$), and then 30 m of heterogeneously textured norite that shows locally pegmatoidal texture, may be veined by felsic material and may contain enclaves or intrusives of fine-grained norite (e.g., at a depth of 888 m). Below the norite is a further 20-m layer of pegmatoidal harzburgite (M1$_L$), olivine pyroxenite, and, locally, pyroxenite containing abundant felsic veins and large (several centimetre-wide) sulfide pods. This is underlain (from ~ 916 m) by a heterogenous sequence of calcsilicate xenoliths within pegmatoidal norite. The borehole was terminated at 925.77 m.

As expected, MgO contents are relatively high in the harzburgites of M1$_L$, but lower in orthopyroxenite of M1$_U$ and norite. Interestingly, both M1$_L$ layers show a broad trend of upward increasing MgO content, analogous to the observation of Yudovskaya et al. (2017a, b) in drill core UMT314. The harzburgites have relatively low Cr/MgO (0–50) due to the predominance of olivine, whereas pyroxenites have higher Cr/MgO of 80 to > 200, with the highest values reflecting the presence of chromite. All rocks of the reef interval (i.e. BAR, M2, M1) contain elevated PGE, at > 400 ppb. There is a broad general trend of a downward decrease in PGE through the profile, an observation that has been made across much of the Turfspruit project area. PGE contents are particularly high in M2 and M1, but in addition, levels are also elevated in the noritic pegmatoid between the M1 layers, notably in some large pods of sulfide. Pt/Pd is mostly around unity, but sharply higher in M2 and the Middling unit, a trend that is found across the project area. Au/Pd broadly mirrors the Pt/Pd trend. Ca/Al$_2$O$_3$ is mostly 0.5–1, overlapping with typically magmatic values found in the UCZ of the WBC. However, higher, putatively non-magmatic values of Ca/
Fig. 4  Compositional variation of selected major and trace element concentrations as well as element ratios in representative drill cores intersecting the Flatreef
Al$_2$O$_3$ are found in M$_{1_U}$, as well as some of the relatively more pyroxene-rich portions of M$_1$ and in the footwall parapyroxenites.

Drillcore UMT341d1 intersected a thick pyroxenitic reef interval, from the Bastard Reef into the footwall of M$_1$, with only thin intervals of olivine-rich rocks. The hanging wall of BAR consists of mottled anorthosite that has a sharp basal contact to sulfide-bearing BAR pyroxenite, which in turn directly overlies M$_2$. The boundary between BAR and M$_2$ is marked by a 2–3-m relatively PGE depleted pyroxenitic interval. M$_2$ is underlain by nearly 60 m of predominantly pegmatoidal orthopyroxenite classified as M$_{1_U}$. There is a 50-cm granite vein in the upper portion of M$_{1_U}$, and a 7-m granite interval at around mid-level. Both granite veins have harzburgite in their footwall and/or hanging wall (Electronic Supplementary Materials (ESM) Fig. 1a-b). There is also a 1.2-m fine- to medium-grained norite in the central portion of M$_{1_U}$ that has sharp contacts to the pyroxenite host rock (ESM Fig. 1c). The base of M$_{1_U}$, i.e. the contact to the footwall pyroxenite, is defined purely on textural grounds, by a change from pegmatoidal to medium-grained pyroxenite. In compositional terms, there is no clear break between these two lithologies.

While MgO contents of the rocks fall into a relatively narrow range of 20–25%, Cr/MgO ratios show more variation than in many other drill cores, due to many samples being enriched in chromite, in the form of thin schlieren, lenses, or stringers, as well as small irregular blebs. Notably, the chromite is commonly set in a matrix of plagioclase, in common with many other Flatreef drill cores. As in drill core UMT263, PGE contents show a broad trend of progressive downward decrease. In addition, PGE contents sharply decrease at a level a few metres above and below the main granite intrusive and around the norite layer/enclave. As in most other Flatreef cores (with the exception of drill core UMT263), Pt/Pd ratios decrease with depth. M$_1$ has markedly lower Pt/Pd (around unity) than M$_2$ and other hanging wall rocks (> 1). Au/Pd again broadly mirrors the Pt/Pd trend. CaO/Al$_2$O$_3$ is relatively homogenous at around 0.5–1 suggesting relatively little input of the country rocks. Sulfide contents are mostly 1–4%, with a few samples containing up to 8% sulfide. The lowest sulfide contents (< 0.5%) are found in the norite and its hostrocks.

Bore hole UMT378 was collared in Zone 1, approximately 1 km to the NW of bore hole UMT263. At the top of the CZ are ~ 5 m of mottled anorthosite (HW1–2). The rocks of the Flatreef show much less lithological variation than in many other drill cores, reflected by a near constant MgO content of 20–40% throughout much of M$_1$, and, with somewhat higher values (up to 1.5) in the basal sample of M$_{1_U}$, the low values reflecting the presence of barren albitic veins. The calcisilicate also has locally highly elevated PGE contents (> 1 ppm). Pt/Pd is at or above unity in M$_2$ and the Middling unit, and slightly lower in much of M$_{1_U}$. Several of the calcisilicate samples have strongly elevated Cu/Ni (not shown), reflecting the presence of relatively Cu-rich sulfides. CaO/Al$_2$O$_3$ is mostly 0.6–1, but with somewhat higher values (up to 1.5) in the basal sample of M$_{1_U}$ and, particularly, the calcisilicate.

In summary, the drill core data show that the examined portion of the Flatreef shares a number of similarities, but also important differences with the interval between the Merensky and Bastard reefs of the WBC. Key similarities include (i) the presence of Cr stringers bounding/underlying the reefs, (ii) the common occurrence of pegmatoid and harzburgite at the base of the reefs, and (iii) the presence of elevated PGE contents in
the footwall of the reefs. Key differences include (i) the much greater thickness of footwall mineralisation in the Flatreef (up to 10 s of metres e.g. in UMT263) relative to the Merensky Reef of the WBC (a few centimetres to rarely a few metres; Barnes and Maier 2002). (ii) The development of several mineralised layers within poorly mineralised pyroxenite and norite in the Flatreef, whereas the Merensky and Bastard reefs in the WBC normally form just 1 layer each. (iii) The occurrence, in the Flatreef, of harzburgite (M1_L) below, within and above orthopyroxenite (M1_U), whereas in the MR of the WBC, harzburgite, where developed, always underlies pyroxenite (Viljoen 1999). (iv) The enhanced thickness of Flatreef (M1) pegmatoid (up to 60 s of metres in UMT341d) compared with the Merensky pegmatoid in the WBC (~ 2 m maximum). (v) Whereas the MR of the WBC typically rests several kilometres above the sedimentary country rocks, the high-grade reefs of the Flatreef (M1 and M2) can rest directly on calcsilicate floor rocks (e.g. in UMT263, UMT378, UMT449). (vi) Pt/Pd in the Flatreef and its floor rocks progressively decrease with depth, whereas in the MR of the WBC and its floor rocks Pt/Pd remains largely constant (Maier et al. 2013).

**Thickness variation of Flatreef units**

In Fig. 5, we have compiled contour maps of the thickness of certain key units across the project area. Of particular note is that BCU, M1_U, and M1_L show complimentary thickness variation, i.e., where BCU is thick, the combined M1 package is thin, and where M1_U is thick, M1_L is thin, and vice versa. In addition, the thicknesses of the M1 units appear to be controlled, at least in part, by the changes in dip of the sequence, with enhanced thickness occurring where dip flattens (Fig. 6).

**Binary variation diagrams**

Binary plots of PGE vs. wt% sulfide show well-defined correlations between Cu and Ni with sulfide, and somewhat weaker correlations between PGE, Au, and Co with sulfide (Fig. 7). The positive correlations likely reflect control of metals by sulfide, but individual reefs define distinct trends. The UG2 reef shows the lowest Pt/sulfide and Pd/sulfide ratios of the analysed rocks, followed by M1_L and the Bastard Reef, and with M1_U and M2 having the highest PGE/S ratios. The Bastard Reef is relatively rich in Au, whereas M1_L is relatively rich in Ni and Co. In terms of Cu/S, all units show considerable overlap. When Pt and Pd are plotted against each other (Fig. 8a), mostly good positive correlations are evident, but with subtle differences for individual units. The Middling and HW units are strongly Pt enriched, BAR and M2 are slightly Pt enriched, whereas M1, UG2, and the FW rocks are variably Pd enriched. Significant scatter is confined to the HW and FW rocks of the reefs. The average Pt/Pd of M2 is higher than that of the MR in the WBC (Table 1). The correlation between Pd and Cu is relatively poor (Fig. 8b). Most samples have Cu/Pd significantly below primitive mantle (~ 7000; Barnes and Maier 1999), with M2 and M1_U having the lowest Cu/Pd, M1_L, UG2 and the Bastard Reef having slightly higher Cu/Pd, whereas several samples from the Middling, HW and FW units (as well as several Bastard Reef samples) have Cu/Pd at or above PM. Nickel (Fig. 8c) shows generally good positive correlations with Cu. Compositional overlap between units is strong, but

![Fig. 5 Diagram showing thickness variation of a Bastard, b M1_U, and c M1_L layers across the Platreef project area (modified from Peters et al. 2016)]
the UG2 and FW rocks tend to have somewhat higher Cu/Ni ratios than the remainder of the samples.

Pd shows a good positive correlation with Ir (Fig. 8d), but Au/Pd is highly variable, showing relatively low values in the UG2, FW, M1 and M2, but higher values in samples from the Middling unit, HW and BAR (Fig. 8e). These trends are essentially mirroring variation in Pt/Pd. Figure 8f illustrates that most Flatreef units have broadly similar Pt/Ir as the Merensky Reef of the western Bushveld, but strongly elevated Pd/Ir. Both ratios are somewhat lower than in most Platreef rocks (not shown). BAR has systematically higher Pt/Ir than the other units, whereas the UG2 has the lowest Pt/Ir and Pd/Ir.

**Discussion: petrogenesis of the Flatreef**

**Composition and emplacement of the Flatreef parent magmas: constraints from Cu contents**

If the Flatreef is the stratigraphic correlative of the uppermost UCZ in the WBC (Grobler et al. 2019), then it would seem reasonable to propose that the parent magma to the Flatreef was a mixture of Bushveld B1 and B2 magmas, as has been modelled for the Merensky Reef of the WBC (Barnes and Maier 2002). However, the Flatreef rocks are too Cu rich to be explained solely by any mixture between B1 and B2 magmas (Fig. 9). The data are equally inconsistent with crystallisation from relatively more fractionated (i.e. enriched in Cu and depleted in Ir) B1 or B2 magmas than those in the rest of the Bushveld because Flatreef pyroxene, plagioclase, and olivine have broadly similar compositions to those in the uppermost CZ rocks elsewhere (Abernethy et al. 2020).

One could propose that the high Cu contents in the Flatreef are due to in situ sulfide melt fractionation after magma emplacement, but this model is rejected in view of the lack of a complementary component of monosulfide solid solution (MSS) in the Flatreef. Alternatively, Cu-rich magmatic sulfide could have been assimilated prior to final emplacement, in a staging chamber, followed by entrainment of the sulfides during continued ascent of the magma (Lee 1996; McDonald and Holwell 2007). This model could potentially explain the vastly larger PGE budget of the Flatreef, and indeed the entire northern limb, compared with the UCZ in the WBC; However, conclusive evidence for sulfide melt saturation of Bushveld magmas prior to final emplacement is not documented. For example, McDonald and Holwell et al. (2007) argued that signatures of metal depletion in some northern lobe LZ rocks reflect sulfide melt saturation prior to emplacement of the LZ, but undepleted LZ sequences also exist (Yudovskaya et al. 2013). Holwell et al. (2011) found sulfide inclusions in Platreef chromite crystals and interpreted both to have been entrained from depth. Our main objection to the sulfide entrainment model is that the fine-grained marginal rocks of the Bushveld Complex, which are generally accepted to represent the parent liquids to the intrusion, are undersaturated in sulfide melt (Barnes et al. 2010). Also, most global
PGE reefs are situated near the transition from ultramafic to mafic rocks (Maier et al. 2015), suggesting that in situ fractionation plays a dominant role in reef formation.

A more plausible model for the relative Cu enrichment of the Flatreef is addition of country rock Cu to the mixed B1/B2 parent magma. The shales of the Duitschland and Malmani formations have average contents of 20–25 ppm and 40 ppm Cu, respectively (Baglow and Brandl 2009). In contrast, dolomite of the Malmani Formation has < 5 ppm Cu and is thus unlikely to be an important source of external Cu. Bearing in
mind that significant in situ assimilation of bulk crust or partial melts of crust is inconsistent with the overlap in silicate mineral composition between the Flatreef and the UCZ elsewhere in the Bushveld (Abernethy et al. 2020). Cu (and S) were likely introduced selectively, i.e., by a fluid that dissolved country rock sulfides.

Hydrodynamic cumulate sorting and sieving during granular flow

Maier et al. (2013) applied a model of crustal loading and subsidence, previously proposed for certain sedimentary basins (Howell and van der Pluijm 1999) to the Bushveld magma chamber. The model is consistent with the lopolithic shape of the Bushveld and many other layered intrusions. The authors further proposed that seismicity associated with subsidence, notably preceding the emplacement of the voluminous Main Zone magma pulse(s), mobilised incompletely consolidated crystal slurries at the top of the Critical Zone crystal pile and caused down-dip slumping of the slurries towards the centre of the Bushveld lopolith. The slurries underwent phase sorting through kinetic sieving resulting in layering, with pyroxenes (+chromite and sulfide) concentrated near the base of flows, and feldspathic cumulates deposited in the upper portion of flows. Apophyses of the UG1 chromitite and the Merensky Reef (Maier and Barnes 2008; Latypov et al. 2017; Mitchell et al. 2019) and exposures of anorthosite seams at Maandagshoek (Maier et al. 2016b) suggest that crystal slurries locally injected into their floor and roof cumulates. The slurry model was tested experimentally by Forien et al. (2015) and is consistent with microstructural studies that show abundant supra-solidus deformation textures.
interpreted to indicate deformation of a crystal-rich mush undergoing shearing (Vukmanovic et al. 2018, 2019). Other features in layered intrusions that are consistent with slurry flow include structures interpreted as cross bedding and soft sediment deformation (Irvine et al. 1998), and the thickening of layers in potholes (Maier et al. 2013).

Grobler et al. (2019) presented drill core evidence in support of hydrodynamic processes in the Flatreef, including slumping structures at various scales (centimetres to 100s of metres), schlieren banding and flow structures, notably in the UG2 chromitite. We highlight here thickened intervals of mineralised pyroxenite that show little evidence of fractionation, as reflected by nearly constant MgO contents and Cr/V ratios across up to 80 m of depth (Fig. 4). We suggest that the thickened intervals formed through accumulation of pyroxenitic crystal mushes in flow dynamic traps, essentially representing large potholes. Other features of the Flatreef consistent with efficient hydrodynamic processes include the relatively low plagioclase contents of Flatreef ultramafic rocks compared with those of the UCZ in the WBC, as well as apophyses of M2 in M1, and of M1 in its footwall (Fig. 4).

Volatiles may also play an important role in hydrodynamic sorting. The solubility of water in basaltic magma is strongly pressure dependent. Thus, in shallow magma chambers such as the Bushveld, emplaced at a depth of 1–3 km (Cawthorn and Davies 1983; Wallmach et al. 1989), it is likely that much of the dissolved H$_2$O in the magma exsolved during intrusion. In the case of magma batches that intruded as sills into a largely consolidated cumulate pile, volatiles could be temporarily trapped, to be released by earthquakes associated with magma emplacement, together with CO$_2$ and H$_2$O formed during devolatisation of, and temporarily trapped within, dolomitic country rocks. In flowing magmas, particularly relatively viscous crystal mushes, gas bubbles may deform and coalesce and act as lubrication planes to substantially decrease the viscosity of the mush (Lesher and Spera 2015). This may facilitate slumping of crystal mushes or wholesale sliding and thrust duplication of semi-consolidated cumulate packages (see Appendix 5 in Grobler et al. (2019)), with the trapped fluids that acted as lubrication planes at the base of the thrust sheet ensuring the lithological integrity of the sheet (King Hubbert and Rubey 1959).

A further potential factor in facilitating crystal sorting could be grain size. Lai et al. (2017) have shown that fine-grained layers can act as lubrication planes for granular flows. This model may be relevant in the case of Cr stringers that formed due to kinetic sieving along the base of ultramafic layers.

**Recrystallisation of the Flatreef**

Fluids or volatile-rich residual melts generated during cooling and crystallisation of cumulates or during metamorphism of footwall rocks may ascend through the crystal pile and locally remelt and recrystallise the rocks, notably at layer contacts (Bédard et al. 1988; Holness et al. 2006; Solano et al. 2014). These concepts were applied to explain the formation of the Merensky Reef by previous researchers (e.g. Ballhaus> and Stumpfl 1986; Boudreau 1988, 2008, 2016b; Boudreau and McCallum 1992; Nicholson and Mathez 1991; Mathez 1995; Mathez and Kinzler 2017). Specifically, it was proposed that ascending hydrous melts and volatiles were trapped at the contact between the Merensky pyroxenite and its footwall norite, both being incompletely solidified. The fluids/hydrous melts lowered the solidus and liquidus temperatures of the rocks and caused an expansion of the olivine and chromite stability fields at the expense of pyroxene and plagioclase (Mathez and Kinzler 2017). In both layers, plagioclase and pyroxene underwent partial melting, the latter also peritectically reacting to olivine. In norite, some plagioclase was preserved, particularly if its proportion in the rock was initially supercotic to form an anorthosite restite (Mathez and Kinzler 2017). The lower and upper contacts of the melt layer with the residual cumulates represent reaction fronts where chromitite stringers crystallised. Where some of the most volatile-rich melt was trapped, peritectic olivine reacted back to orthopyroxene to produce an orthopyroxenite pegmatoid, but where the volatile-rich residual melt escaped, harzburgitic pegmatoid was preserved. The composition of

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**Fig. 9** Ratio diagram of Ni/Pd vs Cu/Ir for key reef intervals in drill cores UMT 314, 336, 341D, 345, 378, and 390. Modelling assumes MSS fractionation of a sulfide melt that segregated from Bushveld B1 magma and a 60:40 mixture of B1 and B2 magma at various R factors. Assumed sulfide melt/silicate melt D values for Pd and Ir are 30,000, for Cu 1000, and for Ni 500. Assumed MSS/sulfide melt D values for Ir is 2.53, Pd 0.15, Cu 0.22, and Ni 1.37 (Barnes et al. 1997). Note that the data trend cannot be explained by a parent magma of B1 composition, but requires addition of significant Cu (in our case 90 ppm)
the proposed melts/fluids remains poorly constrained. In a fluid inclusion study of the Merensky Reef in the WBC, Zhitova et al. (2016) documented the presence of CO$_2$- and CH$_4$-enriched fluids as well as brines.

The model of flux melting and recrystallisation has been highly controversial amongst Bushveld researchers. In part, this is because the cumulates contain few hydrous silicates. The most comprehensive documentation of the modal abundance of mica is that of Teigler and Eales (1996). They reported mostly <1% phlogopite and hornblende in the LZ and lower portion of the CZ, albeit with many samples having several per cent mica. Similarly, Karykowski and Maier (2017) showed that the LZ can locally be highly enriched in phlogopite (up to 6 modal %). Maier and Eales (1997) observed that within the UCZ, olivine-bearing cumulates tend to be relatively mica rich, and Maier (1995) reported abundant small (<0.5 mm) phlogopite from an olivine-bearing UCZ troctolite. These data suggest that the proportion of mica could have been underestimated in Bushveld cumulates. Also, Boudreau (2019) argued that vapour saturation may occur at a temperature beyond the stability of phlogopite, suggesting that scarcity of mica is not a reliable measure for the timing of vapour saturation in cumulates.

Another reason why the recrystallisation model remained controversial is that the Merensky pegmatoidal pyroxenite has been shown to have similar incompatible trace element contents and ratios to other, non-pegmatoidal pyroxenites in the UCZ (Barnes and Maier 2002). However, it could be argued that melt generated by fluid fluxing caused increased porosity that facilitated escape of the hydrous melt, thereby resulting in relatively low contents of mica and incompatible trace elements. Furthermore, Li et al. (2005) and Boudreau (2016a) reported that in the Merensky and J-M reefs, melt inclusions within chromite, olivine, and apatite are composed largely of biotite and Na-amphibole and lesser amounts of olivine and pyroxene. These alkaline inclusions may represent the residual/partial melt.

Cumulate recrystallisation triggered by volatile-rich melts or fluids would provide an explanation for several other observations in the Bushveld Complex and other layered intrusions, including (i) pervasive reversed zonation in plagioclase of the UCZ (Maier 1995; Maier and Eales 1997; Maier et al. 2016b; Hunt et al. 2018) and in the Stillwater Complex (Czamanske and Schéidile 1985; Boudreau 1988; Baker and Boudreau 2019), (ii) elevated Sr ratios below and within certain ultramafic layers (UCZ chromitite-Kinnaird et al. 2002; UG2-Schannon et al. 2018), and (iii) the occurrence of LG6 chromitite with unusually high Cr/Fe and hosted by unusually olivine-rich silicate rocks, in pothole-like depressions at the Ruighoek mine (Latypov et al. 2018) which we interpret to have resulted from channelised fluid flow triggering oxidation of Fe in chromite and recrystallisation of pyroxenite to harzburgite.

The recrystallisation model was first applied to the Platreef by de Waal (1977) and to the Flatreef by Marquis (2015) to explain, amongst other features, the pegmatoidal textures and abundance of olivine, notably in M1$_L$. The model offers a possible explanation for the observed variation in relative thickness of orthopyroxenite and harzburgite across the property (Fig. 5)—where orthopyroxenite is thick, harzburgite is thin, and vice versa—in that both rock types could be reaction products of an originally medium-grained orthopyroxenite-harzburgite interval. The model would potentially also provide an explanation for several other observations namely (i) the preferred occurrence of M1$_L$ in areas where the dip of the sequence changes significantly which could be interpreted to favour early-stage trapping of fluids and volatile-rich melts percolating upwards along layer contacts (Fig. 6), (ii) the subtle depletion in incompatible trace elements in M1$_L$ relative to M1$_U$ (Table 1) that could reflect late-stage escape of residual volatile-rich melt thereby stabilising olivine, and (iii) the association of chromite with intercumulus plagioclase in many Flatreef chromitite stringers and seams (Fig. 10) that could result from fluid-driven partial melting and recrystallisation of pyroxenite (Boudreau 1988; Nicholson and Mathez 1991).

The origin of the Cr stringers has remained particularly controversial since Lee et al. (1983) proposed a secondary origin for Cr stringers enveloping pyroxenite aggregates of the “Boulder Bed” and bracketing the Merensky pegmatoid. Analogous chromite envelopes have been described in Flatreef pyroxenite lenses (Grobler et al. 2019). Chromite grains in the Merensky Cr stringers have highly irregular, amoeboidal textures previously explained by annealing (Eales and Reynolds 1986), corrosion (Cawthorn and Boerst 2006; Hutchinson et al. 2015), or supercooling (Vukmanovic et al. 2013). Scoon and Costin (2018) reported that the stringers can be enriched in phlogopite and rutile, and that plagioclase in the stringers tends to be reversed zoned and has higher An content than plagioclase in the adjacent hostrocks, with some values exceeding An$_{90}$. Some of these observations have also been made for the Lone Cr stringer within the footwall of the Merensky Reef of the WBC (Maier and Eales 1997). Boudreau (1988) and Nicholson and Mathez (1990) proposed that the Cr stringers crystallised from a volatile-rich melt, consistent with the presence of mica in some stringers. In contrast, Scoon and Costin (2018) argued that the Cr stringers formed through partial melting of gabbronite or anorthosite caused by late-stage intrusion of ultramafic sills, with chromite precipitating from the hybrid contact melt (cf O’Driscoll et al. 2010). However, some Bushveld Cr stringers such as the Lone Cr stringer (Maier and Eales 1997) occur within anorthosite or norite where a potential heat source that might have triggered melting is not apparent.

Kennedy et al. (2018) have proposed a recrystallisation model for the Troctolite Unit in the Bushveld northern lobe.
It is possible that other olivine-rich lithologies in the northern lobe and elsewhere in the Bushveld formed in an analogous manner. For example, in the WBC, the so-called “Pseudoreefs” consist of coarse-grained pegmatoidal troctolites or harzburgites with large subhedral or anhedral olivine grains set in a matrix of interstitial plagioclase and/or orthopyroxene (Eales et al. 1988; Maier and Eales 1997) resembling M1L in the Flatreef. Similarly, the PGE reefs of the Waterberg project in the Bushveld northern lobe (Kinnaird et al. 2017) are hosted in relatively coarse-grained troctolites, with olivine predominantly associated with interstitial plagioclase, and strongly recrystallised sulfides showing disequilibrium textures, lobate grain boundaries, and alteration with abundant magnetite veins.

One unresolved aspect of the recrystallisation model is that few obvious melt/fluid channels are evident. The Merensky Reef may contain sub-vertical plagioclase-rich veins (Barnes and Maier 2002), but in general these are rare. Our interpretation is that hydrous melt could have migrated along inclined bedding contacts (Fig. 11) to end up in the peripheral portions of the complex that are now eroded. Bedding-parallel flux of volatile-rich melt or fluids and resulting autometasomatism within layers could explain the sharp changes in mobile ITE contents (Rb, K, LREE, C) between adjacent layers and the upward increase in these elements within layers, resulting from trapping at the upper contacts of layers (Fig. 12).

Mobility of sulfide during cumulate recrystallisation

In both the Flatreef and the Merensky Reef of the WBC, the PGE mineralisation commonly extends into the floor rocks of the reefs (Fig. 13). However, in the WBC, sulfides are typically confined to the uppermost few centimetres to decimetres of the footwall (Barnes and Maier 2002), whereas in the Flatreef, the mineralisation extends for up to several 10 s of metres into the floor (UTM231, 232, 263, and 345). Furthermore, in the case of the Flatreef, PGE also occur within calcilicate floor rocks (UTM449, 421, and 417). In the WBC, the Merensky footwall mineralisation is normally interpreted to have resulted from late-magmatic percolation of sulfide melt (Leeb-du Toit 1986; Barnes and Maier 2002; Roberts et al. 2007; Naldrett et al. 2009). The more pervasive nature of footwall mineralisation in the Flatreef could thus suggest that late-magmatic sulfide mobility played a relatively more important role. The model is consistent with textural evidence from micro XRF (Fig. 3), suggesting that sulfides are strongly interconnected in 3D and represent networks of sulfide liquid percolating down, and displacing silicate liquid percolating up (Barnes et al. 2018). Sulfide liquid percolation is also suggested by enrichment of PGE in some of the normally barren calcilicate intervals (Abernethy et al. 2020). Relatively high Pd/Ir and Cu/Ni in some calcilicate samples (Abernethy et al. 2020) indicate PGE and base metal fractionation during transport. Of further importance is the observation that PGE grades decrease in a wide (up to 5 m) halo around a norite vein within the M1 pyroxenite of UMT341D1 (Fig. 3). Our interpretation is that norite intruded incompletely solidified pyroxenite temporarily increasing the porosity of the rock and leading to percolation of sulfide in a down-dip, layer-parallel direction, resulting in metal poor pyroxenite.

Potential explanations for the decreasing Pt/Pd with depth in the Flatreef

The Flatreef and its footwall mineralisation is typically characterised by a broadly progressive downward decrease in Pt/Pd, from values as high as 2 in BAR and M2 (and locally the UG2), to values around or just below unity in M1, and as low as < 0.5 in the footwall assimilation zone (Grobler et al. 2019; Fig. 4 of present study). A similar downward decrease in Pt/Pd was documented in many other magmatic ore deposits, including the Finnish PGE-rich layered intrusions (Iljina et al. 2015), Duluth (Thériault et al. 2000), Raglan/Cape Smith (Barnes et al. 1992), and Muskox (Barnes and Francis 1995). The trends cannot be explained by sulfide percolation and associated
Fig. 11  a Schematic model of layer formation in Bushveld UCZ, including chromitites and PGE reefs. Note presence of several fluid-migration directions: crystal slurries move downwards along top of crystal pile forming ultramafic layers (including chromitites). Feldspathic slurries move upwards, forming anorthosite (yellow arrows). Sulfides locally percolate downward from ultramafic slurries (red arrows). Hydrous fluids percolate up-dip along layer contacts (pink arrows). Harzburgites may form either in laterally moving metasomatic fronts along the compacted base of ultramafic layers (b), or through intra-sill reactive flow (c).

Fig. 12  Variation in concentration of a MgO, b carbon, and c La/Yb in drill core UMT314. Note enrichment in C and La/Yb within the M1 pegmatoids.
MSS crystallisation because Pt and Pd have similar D values with regard to MSS (Fleet et al. 1993).

(i) Mobility of Pd in fluids

Empirical (Barnes and Liu 2012; Holwell et al. 2017) and experimental data (Hanley 2005) indicate that at typically magmatic conditions (sulfide present, $f_O^2$ near QFM, and near-neutral pH) Pd is more mobile in fluids than Pt. Based on these data, one could suggest that any fluids migrating through the basal portions of layered intrusions were enriched in Pd relative to Pt (Boudreau 2019). Palladium could precipitate in sulfide-rich rocks, potentially offering an explanation for the relatively high Pd/Ir and Pd/Pt of the lower Flatreef (Fig. 8f). However, the Flatreef rocks show good correlations between individual PGE in binary variation diagrams (Figs. 7 and 8) providing little evidence for significant selective PGE mobility.

(ii) Variable partitioning of PGE into sulfide

Several experimental studies have shown higher partitioning of Pd than Pt into sulfide melt (Barnes and Lightfoot 2005 and references therein). This could result in the residual melts becoming relatively Pt enriched, resulting in progressively more Pt-rich sulfides with height (Fig. 4).

(iii) Pt depletion of earliest Flatreef magma pulses

The earliest Flatreef pulses could have been relatively Pt depleted due to Pt alloy fractionation prior to final emplacement, in response to contamination with reduced (graphite-bearing) sedimentary rocks of the Duitschland Formation. Borisov and Palme (1997) have shown that Pt solubility is positively correlated to oxygen fugacity.

Summary and conclusions

The growing body of information including the new data presented in this paper suggests that the PGE-Ni-Cu ore deposits of the Flatreef formed through a combination of magmatic, hydrodynamic and hydromagmatic processes, starting with (i) the influx of several batches of PGE fertile basaltic magma. The magmas became saturated in an immiscible sulfide melt due to assimilation of country rock sulfides during emplacement resulting in somewhat higher Cu contents of the Flatreef than in the reefs of the remainder of the Bushveld.
Assimilation was particularly pronounced in the initial magma batches from which the lower portion of the Flatreef crystallised. Sulfides in these rocks are relatively Pd enriched, possibly due to higher $D_{\text{sulfide melt:silicate melt}}$ of Pd than Pt, mobility of Pd in late- and post-magmatic fluids or Pt depletion of the earliest magma pulses resulting from Pt alloy fractionation triggered by contamination with reducing country rocks. We do not subscribe to the model of entainment of PGE-rich sulfides from a staging chamber, mainly because all of the fine-grained marginal rocks to the intrusion that are generally considered to represent the parent magmas to the cumulates are sulfide undersaturated. (ii) Deposition of a thick pile of mafic-ultramafic cumulates caused subsidence of the magma chamber. Aided by flux of volatiles from the crystallising cumulates and the devolatising country rocks, subsidence and associated seismicity facilitated slumping of crystal slurries located at the top of the growing cumulate pile towards the centre of the Bushveld lopolith and into local troughs. This resulted in sorting and compaction of the slurries, pronounced layering and localised hydrodynamic erosion of floor cumulates (Fig. 13). (iii) Contacts between compositionally and texturally distinct layers acted as traps for volatiles that ascended through the cumulate pile triggering partial melting and recrystallisation of cumulates to form pegmatoidal rocks that may be highly enriched in sulfides, secondary olivine and, in some cases, graphite and phlogopite. (iv) The partial melting of the Flatreef cumulates facilitated downward percolation of sulfides, locally for up to several 10 s of metres. In places, ultramafic cumulates injected into their fluidised, volatile-rich footwall rocks forming extensive sill-like apophyses (Fig. 13). The combination of these processes resulted in pervasively recrystallised (zone refined) cumulate packages hosting exceptionally thick PGE mineralised intervals (up to more than 300 m at > 1 ppm PGE).

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