Title: Quantifying chemical weathering intensity and trace element release from two contrasting basalt profiles, Deccan Traps, India

Keywords: Deccan Traps, weathering index, mafic index of alteration, index of lateritisation, Sm/Nd, europium anomaly, dust

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Abstract: Weathering profiles developed on basalt substrate contain information relevant to climate, atmospheric composition and evolution, nutrient release into the hydrosphere, and understanding Martian regolith. In this study, the chemical compositions of two profiles developed on Deccan Trap basalt are examined. One is sub-Recent and has only progressed to a moderate degree of alteration (Chhindwara profile), whereas the other is ancient (Paleocene) and the degree of alteration is extreme (Bidar laterite). In an attempt to better quantify the chemical changes during incipient to intermediate weathering of mafic substrates, we propose a new index: the mafic index of alteration (MIA). Similar to the chemical index of alteration (CIA), the MIA quantifies the net loss of the mobile major elements (Ca, Mg, Na, K ± Fe) relative to the immobile major elements (Al ± Fe). The redox-dependent weathering behaviour of Fe is factored into two separate arrangements of the MIA that apply to oxidative [MIA(O)] or reduced [MIA(R)] weathering. The MIA can be visualised in a variety of ternary diagrams in the Al-Fe-Mg-Ca-Na-K system. To chemically quantify the stages of advanced to extreme weathering, at which the MIA and CIA are ineffective, the SiO2 to (Al2O3+Fe2O3) mass ratio, based on the established Si-Al-Fe (SAF) 'laterite' ternary diagram, is used; we propose that this ratio be referred to as the 'index of lateritisation' (IOL).

Major element chemical variations, as expressed by weathering indices, were used to relate the extent of weathering with the behavior of trace elements (alkali, alkaline earth, rare earth, and Nb) in the profiles. During the early stages of basalt weathering, the mobile trace elements (Sr, Be, Li) are anti-correlated with the chemical weathering indices and thus released during these stages. By contrast, the monovalent elements (K, Rb, Cs, Tl), excluding Na and Li, appear to be associated with the pedogenetic clay minerals. Of these elements, those with the closest ionic radii are most closely related. Fractionation of the REE (Sm/Nd, Eu/Eu*, Ce/Ce*) is evident during weathering of the basalt. The loss of Eu is linked with Sr, Ca, and Na and thus plagioclase dissolution during the stages of incipient to intermediate weathering. The fractionation of Sm/Nd suggests that basaltic weathering products may not always preserve their parent rock ratio and, consequently, their Nd isotope composition over time.

Finally, weathering in the sub-Recent profile is shown to have progressed across two lava flows, whose morphology initially controlled the extent of weathering. Certain compositional variations in the
original flows (e.g., immobile element ratios) are preserved through the effects of chemical weathering and have the potential to influence mass balance calculations across the entire profile.
CHEMGE6928 highlights

- New chemical weathering index is introduced: the mafic index of alteration (MIA)
- Si:(Si+Fe+Al) ratio to quantify advanced weathering (the index of lateritisation)
- Separate lava flows in a weathering profile influence mass balance calculations
- REE fractionation during basalt weathering: LREE/HREE, Sm/Nd, Ce/Ce*, Eu/Eu*
Quantifying chemical weathering intensity and trace element release from two contrasting basalt profiles, Deccan Traps, India

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ABSTRACT
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HIGHLIGHTS

- New chemical weathering index is introduced: the mafic index of alteration (MIA)
- Si:(Si+Fe+Al) ratio to quantify advanced weathering (the index of lateritisation)
- Separate lava flows in a weathering profile influence mass balance calculations
- REE fractionation during basalt weathering: LREE/HREE, Sm/Nd, Ce/Ce*, Eu/Eu*
1. INTRODUCTION

Chemical breakdown of thermodynamically unstable minerals at the Earth’s surface by weathering agents is a fundamental part of global elemental cycles. It is responsible for the long-term regulation of global atmospheric CO₂ concentrations and for generating clastic sediment and dissolved elements for delivery to fluvial and marine reservoirs. Therefore, in view of this critical role, it is perhaps not surprising that several geochemical indices have been introduced for the purpose of understanding and quantifying the processes of weathering (e.g., Parker, 1970; Kronberg and Nesbitt, 1981; Nesbitt and Young, 1982; Harnois, 1988; Maynard, 1992; Fedo et al., 1995; Retallack, 2001 and references therein; Ohta and Arai, 2007; Sheldon and Tabor, 2009; Nordt and Driese, 2010). Such quantification of the degree of weathering in soils/paleosols or clastic sedimentary sequences has proven important from the scale of an individual weathering profile up to that of global mass balance. At the scale of an individual weathering profile, the most useful weathering indices are those with elemental components that can be determined routinely and examined graphically to provide further insight into the weathering and/or post-weathering processes. One of the most widely used examples is the chemical index of alteration (CIA: Nesbitt and Young, 1982, 1984, 1989) and the accompanying A–CN–K diagram (Nesbitt and Young, 1984; Fedo et al., 1995).

In this study, a new chemical weathering index modelled after the CIA is introduced: the mafic index of alteration (MIA). The MIA adds the elements Fe and Mg to the Al–Ca–Na–K system, allowing the mafic mineral component of rock weathering to be quantified. The MIA is most suitable for, but not restricted to, studying mafic rock weathering. The study of chemical weathering of mafic substrates, compared to felsic substrates, provides many advantages from a geological and chemical perspective: a) the parent rock is often fine-grained and typically quite
homogeneous from a chemical and mineralogical perspective; b) the substrate is quite rich in redox-sensitive elements, such as the transition metals; c) basalt or its metamorphosed/altered equivalent is ubiquitous throughout the known geological record, and e) it is of increasing importance for interpreting Martian substrate alteration. Silicate weathering is perhaps the most important removal mechanism of CO$_2$ from the atmosphere, with the weathering of mafic substrates often being cited as especially important in the regulation of global climatic changes (Louvat and Allègre, 1997; Schwarz, 1997; Gaillardet et al., 1999; Taylor and Lasaga, 1999; Dessert et al., 2001; Das et al., 2005; Navarre-Sitchler and Brantley, 2007).

Although the major elements closely reflect the mineralogical transformations during weathering, the cycling of certain minor and trace elements has potential to provide further insight into pedogenetic processes, such as biological activity or the oxidation state of a profile. Linking the release or retention of these elements to specific stages of chemical weathering can be accomplished through the combined use of major element weathering indices and trace element mass balance. In this study, this is demonstrated for the alkali, alkaline earth, and rare earth elements during different stages of alteration of the Deccan Traps basalt.

2. BASALT CHEMICAL WEATHERING AND LATERITISATION

2.1. Incipient to intermediate weathering

For typical basaltic rocks, the susceptibility of minerals to chemical weathering follows the order of glass $\approx$ olivine $>$ plagioclase $\approx$ pyroxene $>$ Fe-Ti oxide (e.g., Eggleton et al., 1987; Nesbitt and Wilson, 1992), although some exceptions related to particular textural characteristics or the weathering environment have been reported (e.g., Craig and Loughnan, 1964). Ultimately,
pedogenetic phyllosilicates (e.g., kaolinite and the smectite group minerals) and Fe and Mn oxides or oxyhydroxides form from the primary basaltic constituents during weathering. The exact pedogenetic mineralogy and order of mineral formation depend on the parent rock composition, climate, biology, redox state, and drainage of the profile (e.g., Prudêncio et al., 2002; Rasmussen et al., 2010). During the incipient to early stages of basalt weathering, 2:1 layer phyllosilicates typically form and significant hydration occurs. As weathering progresses from to intermediate and advanced stages, 1:1 layer clays such as kaolinite or halloysite are typical stable end-products. Therefore, this realm of chemical weathering is sometimes referred to as ‘kaolinitisation’.

During the early stages of basalt weathering, a general net loss of the mobile elements (Mg, Ca, Na, ± K) accompanies the mineralogical transformations (e.g., Nesbitt et al., 1980; Chesworth et al., 1981; Kronberg and Nesbitt, 1981). By contrast, Al, Fe, and Si are predominantly retained. Therefore, quantification of weathering intensity during these stages typically assumes conservation of Al and measures the loss of the mobile elements (Ca, Na, K, Mg). The weathering behaviour of Fe, however, may be more complicated and dependent on the redox and drainage conditions of the profile (e.g., Driese, 2004).

2.2. Advanced to extreme weathering and lateritisation

The term ‘laterite’ comes with a history of debate and disagreement related to the origin of duricrusts (for recent discussion see Ollier and Galloway, 1990; Bourman, 1993; Bourman and Ollier 2002, 2003; Schellmann, 2003). We prefer to adopt the genetic distinction between ferricrete and laterite (e.g., Aleva, 1994; Widdowson, 2007); in this study, ‘lateritisation’ will refer to the progressive, in situ alteration of rock. By contrast, ferricretes are alteration profiles
that are largely generated by the allochthonous input and accumulation of ferruginous cement within existing subaerial substrates. In practice, the distinction between laterite and ferricrete cannot always be readily made in the field, and may require very detailed characterisation of duricrusts (e.g., Beauvais, 1999). Genetic distinctions are further complicated on a smaller scale since even in situ weathering profiles are subject to some lateral element transfer and the incorporation of aeolian sediment (Figure 1).

Lateritisation is generally associated with intense and prolonged leaching conditions in sufficiently hot and humid climates. During advanced to extreme weathering that is associated with lateritisation, the stage of pedogenetic phyllosilicate accumulation is exceeded and Fe- and Al-oxyhydroxides and sesquioxides (e.g., gibbsite, goethite, hematite) begin to dominate the weathering profile mineralogy. The Si largely retained in the phyllosilicates during earlier stages of weathering is lost (e.g., Widdowson and Gunnell, 1999; Hill et al., 2000) and desilication becomes the dominant process modifying the chemistry of the weathering profile. In this respect, the classic chemical definition of ‘laterite’ by Schellmann (1981, 1982, 1986) based on the $\text{SiO}_2$ to $(\text{Fe}_2\text{O}_3 + \text{Al}_2\text{O}_3)$ ratio becomes useful in accompanying field observations of lateritic profiles. This ratio and the accompanying $\text{SiO}_2$–$\text{Al}_2\text{O}_3$–$\text{Fe}_2\text{O}_3$ (SAF) ternary plot provide a quantifiable chemical criterion to examine highly altered substrates, although they are not applicable to allochthonous duricrusts (i.e., ferricretes). The full progression of chemical weathering must be recorded within a single ‘laterite profile’ that is characterised by a gradual alteration of parent rock to saprolite near the base and grading into a pallid zone, followed by a mottled zone, and finally an indurated, Fe-rich duricrust at the top (i.e., the ‘textbook’ laterite profile). Caution should be exercised where an in situ progressive weathering profile cannot be unequivocally identified or where the parent rock is unavailable or unknown.
3. GEOLOGY

3.1 Geological and geochemical context of the Deccan Volcanic Province

The Deccan Volcanic Province (DVP), located in western peninsular India, is the remnant of an extensive, continental flood basalt emplaced during a tholeiitic eruptive acme between ca. 67-65 Ma (e.g., Duncan and Pyle, 1988; Gallet et al., 1989; Widdowson et al., 2000; Hofmann et al., 2000; Chenet et al., 2007; Hooper et al., 2010) at the Cretaceous-Paleogene Boundary (K PgB).

Immediately following the eruption of the Deccan Traps flood basalt, the neo-formed lava fields were exposed to intense tropical weathering due to the equatorial location of the DVP around the KPgB. Accordingly, deep (lateritised) weathering profiles developed uninterrupted throughout the Paleocene. The weathering intensity is likely to have waned in the Eocene following continental uplift resulting from the collision of the Indian and Asian plates and/or denudational uplift of the rifted margin of Western India (65-50 Ma; Beck et al., 1995; Widdowson, 1997); this, together with climate changes resulting from the closure of the Tethyan seaway and later uplift of the Himalayas, and associated profound alteration of Indian continental drainage patterns, are likely to have resulted in changes in Indian weathering regimes (e.g., Kerrick and Caldeira, 1993).

Remnants of Paleocene and Miocene duricrusts are found located at topographical highs in the Western Ghats and across the low-lying Konkan coastal plain, respectively (Widdowson and Cox, 1996; Widdowson, 1997). The high-level duricrusts are interpreted as remnants of the once extensive weathering residuum that had developed on the top of the DVP lava flow package after eruptions had ceased (ca. 60 Ma), whilst the low-lying examples of the Konkan plain represent
alteration of a piedmont-like paleosurface developed at the foot of the Western Ghats escarpment in the upper Tertiary (e.g., Widdowson and Gunnell, 1990; Widdowson, 2007).

Differential weathering and erosion of the flat-lying basaltic flows (c. 4-10 m thick) has produced spurs, ridges and mesas that are separated by flat-floored valleys often containing meandering rivers. East of the Ghats escarpment these rivers drain eastward and southward eventually discharging into the Bay of Bengal, while those that drain the escarpment and Konkan plain flow westward into the Arabian Sea. Increased aridity during the LGM resulted in reduced stream flow magnitudes in the rivers of south and SE Asia, whilst the glacially-induced eustatic low extended river courses across exposed coastal plains. As wetter (monsoon) conditions returned in the early Holocene, the rivers in peninsular India responded by deepening and enlarging their channels to accommodate increased discharge leading to widespread erosion and development of river terraces (e.g., Clift et al., 2002, Kale, 2002; Figure 2). In most areas, these fluvial processes stripped the deeply lateritised Paleocene duricrust and exposed fresh basalt in topographical lows to Holocene weathering. Therefore, most of the current outcrop of the DVP (> 90%) consists of basalt variably weathered to 1 to 10 m in a stepped mesa-like topography that is largely the result of Quaternary (Pleistocene-Holocene) fluvial processes of valley incision and widening in the wake of the last glacial maximum (LGM).

The DVP provides an ideal natural laboratory to study the various stages of chemical weathering of compositionally similar tholeiitic basalt. The thick, near horizontal geometry of the DVP flows and the mesa topography of the area has ensured that the current chemical state of the weathering profiles has been isolated from interaction with ground or surface waters. Further, given the massive areal extent of the DVP flows, the study sites are remote from other basement lithology and only minor aeolian input could possibly interfere with the DVP’s excellent
potential for investigating basaltic chemical weathering (Kisakürek et al., 2004; Wimpenny et al., 2007).

In the present study, two weathering profiles from the eastern DVP, representing very different stages of basalt alteration, are examined. The first is an ancient (Paleocene), highly advanced (lateritic) weathering profile developed in the Bidar area (BB sample series). The second is a post-Pleistocene (sub-Recent) weathering profile located in a roadside quarry near Chhindwara (ChQ sample series) in the Madhya Pradesh district. This profile is substantially less advanced in its degree of alteration, lacking any duricrust development. The geology and sampling strategies of these two chemically different weathering profiles are summarised next and illustrated in Figure 3.

3.2. Bidar profile

The Bidar laterite profile is well exposed at a hillside edge of a mesa near Bidar, Madhya Pradesh, India (17°54.87’ N, 77°32.39’ E; Figure 2). This site is where Newbold (1844, 1846) first suggested laterite to be the consequence of \textit{in situ} rock weathering. The Bidar laterite, sampled in 1998, is interpreted as a deep (~50 m) weathering profile with upwardly increasing alteration and concomitant Fe-enrichment (Borger and Widdowson, 2001; Kisakürek et al., 2004). Between the top and bottom of the profile there is a typical lateritic weathering progression, including a mottled zone, and an indurated duricrust cap (Figure 3a). The unweathered basalt (at a profile depth of > 40 m) gradationally gives way to a corestone-rich horizon and saprolite matrix between a depth of 35-30 m. This depth range is characterised by the typical depletion of mobile alkaline and alkaline earth elements that is associated with incipient weathering. Between 30-25 m, saprolite with Fe-rich mottles and segregations becomes
dominant. Above this point, primary silicates have been completely removed from the profile and Si and Al concentrations decrease relative to Fe. This is associated with an increase in the abundance of irregular Fe-rich agglomerations and a reddening of the saprolite colour upward between ~25-15 m. An unexpected return to bluish-grey saprolite between ~15-10 m marks the only apparent discontinuity in the upward weathering progression. Based on higher than expected Fe concentrations and trace metal accumulation, Kisakürek et al. (2004) argued that this horizon represents the position of a stable paleo-water table. At the highest, and most advanced part of the weathering profile, tabular, semi-indurated laterite at ~10-4 m grades into an indurated laterite cap (4 m and up). Further details of the profile geology are described in Borger and Widdowson (2001) and Widdowson (2007).

Nine samples (BB1-BB9) were collected from the Bidar laterite for previous studies (Mason et al., 2000; Kisakürek et al., 2004; Wimpenny et al., 2007), each representative of zones of key textural change. The focus during the sampling was on the uppermost transition from saprolite to laterite and on representing the various mineralogical and textural changes in the upper, most highly weathered part of the profile. Approximately 1-3 kg of material was taken for each sample to minimise the vertical and horizontal heterogeneity that is common in deeply weathered laterite profiles. The same samples were used for the present study (Figure 3a).

3.3. Chhindwara profile

The Chhindwara District is situated on a 600-750 m elevation basaltic plateau that extends approximately 200 km NE from the Seoni-Chhindwara area toward Jabalpur (Figure 2). Throughout the area, terraced hillsides or low mesas of 50-100 m elevation are separated by flat-floored valleys containing ephemeral rivers draining eastward and southward. The higher
elevation duricrust cap common to the Bidar area is absent. A transect through the Quaternary weathering front, located in a recently abandoned road stone quarry, 7 km east of Chhindwara (22° 04.213’ N; 79° 01.393’ E; Figure 2) was sampled in 2009; see field photographs (Figure 4).

Located at 741 m elevation, the weathering profile extends to a total exposed depth of >6 m across two clearly identifiable massive lava flows exposed in the face of the quarry (Figure 4a). Stratigraphically and geochemically, the two flows appear to belong to the Poladpur-Amenali formation transition zone (Mitchell and Widdowson, 1991). In general, the flow-banded and higher vesicularity areas of both flows are more altered than the thicker, massive regions. This implies that the primary flow morphology has had a strong control on fluid penetration, chemical attack, and, consequently, chemical weathering intensity.

The fully exposed (~4 m) upper flow (ChQB) has weathered to a rust red colour with spheroidal weathering features present throughout and is capped by a thin (~20 cm) soil horizon at the present day surface (Figure 4). The more massive, homogeneous basalt that constitutes most of the flow grades into a zone of weak ‘banding’ at the base (lower 80 cm), which may relate to rheological effects during emplacement and/or minor mineralogical variation. The weathering intensity in the flow, as indicated by the clay content, follows the aforementioned morphology transition, with a sharp increase in phyllosilicates near the base that is coincident with the appearance of the banded texture.

The transition from the upper to lower flow unit is clearly identifiable by the grayish colour of the latter. The colour is presumably a result of the higher abundance of pedogenetic clays and zeolite minerals. In this flow, vesicularity, made evident from zeolite-infilled amygdules, increases towards the flow top and terminates with a brecciated and highly vesicular flow top (~
The base of the lower flow is not exposed beyond the quarry floor. The colour and clay mineralogy are relatively consistent throughout the exposed portion (215 cm) of the lower flow, and suggests a higher (relative to the upper flow) but relatively consistent intensity of weathering. The difference in colour (from the rust red to brown to the greenish-gray) between the upper and lower flows could also indicate that the latter records an earlier weathering history. That is, it may represent a bole bed that formed during a period of volcanic quiescence in the DVP emplacement history (Ghosh et al., 2006; Sayyed and Hundekari, 2006) prior to the eruption of the upper flow. In addition, the presence of zeolite minerals in Deccan basalt is thought to have some control on the chemical weathering progression by retaining elements and inhibiting complete kaolinitisation (Bhattacharyya et al., 1999).

From the Chhindwara profile, a total of 27 samples were collected. The upper flow (ChQB1-B12) and lower flow (ChQA1-A12) are represented by 12 samples each and the remaining samples are from a horizontal profile through a corestone at a depth of 90 cm from the upper flow surface (ChQB9a,b,c,d; Figure 4b). The least-weathered samples of the profile, ChQB12 and ChQB9d, are located in the upper flow at a depth of 140 cm and within the corestone centre, respectively. The sample locations are indicated on a schematic log (Figure 3).

4. METHODOLOGY AND ANALYSIS

4.1. Analytical details

Samples for the Chhindwara profile were pulverised in an agate mill to minimise metal contamination, whereas the powders for the Bidar profile were prepared in a tungsten carbide mill. The major element composition for both profiles was determined by X-ray fluorescence
(XRF) following loss on ignition (LOI) measurement; the Bidar XRF data were obtained at the Open University (UK) and reported previously (Kisakürek et al., 2004; Widdowson, 2007), while the Chhindwara samples were analysed at the Geoscience Laboratories (Sudbury, Ontario). Ferrous iron measurements were also obtained on Chhindwara samples at the Geoscience Laboratories via potentiometric titration with potassium permanganate.

All trace element data were obtained in the Department of Earth Sciences at Laurentian University. A 100 mg powder aliquot was digested in thoroughly cleaned 29 mL PFA screw-top beakers using a 2.5 ml HF plus 0.5 ml HNO₃ acid mixture. After a 72 h digestion period at 160ºC, samples were dried down at 110ºC to drive off SiF₄. The fluoride residue was attacked twice with 0.5 mL 6 N HCl to reduce organic components before double conversion with HNO₃. The converted residue was taken up in 10 g of 20% HNO₃ to yield a nominal 1:100 parts total dissolved solids stock solution. This was transferred to a clear polystyrene test tube and centrifuged to inspect for insoluble fluorides, which were not encountered.

For the trace element analysis, 0.2 g (i.e., 2%) of the stock solution was gravimetrically diluted to 6 g 2% HNO₃ together with a mixture of internal standards (10.6 ppb ⁶Li, 4.4 ppb each Rh, Re and Bi and 1.5 ppb ²³⁵U). A total of 46 trace elements were analysed by quadrupole ICP-MS (Thermo XSeriesII) using the methodology of Eggins et al. (1997) with some modifications (Kamber et al., 2003; Kamber, 2009). Calibration of analyses was performed using the USGS standard W-2 digested under the same conditions as the experiment unknowns using the preferred values reported previously (e.g., Babechuk et al., 2010). Long-term reproducibilities for most elements with this method are between 1-2% (e.g., Kamber, 2009; Babechuk et al., 2010; Marx and Kamber, 2010). The major element and trace element data set relevant to the present study is reported for the Bidar profile in Table 1 and the Chhindwara profile in Table 2.
5. RESULTS AND DISCUSSION – PART I

Major elements, loss on ignition, and quantification of the extent of weathering

In the following sections, the bulk major element trends of the Bidar and Chhindwara weathering profiles are interpreted using the widely recognised chemical index of alteration (Section 5.1), as well as the new mafic index of alteration and index of lateritisation (Sections 5.2 and 5.3). A useable spreadsheet that demonstrates the weathering index calculations can be found in the supplementary materials. Each weathering index is also discussed on accompanying ternary diagrams that allow for the graphical interpretation of the proportional chemical changes.

Included in ternary plots for comparison is a data compilation of published sub-Recent basaltic weathering profiles and Tertiary basaltic laterite and bauxite profiles (Chesworth et al., 1981; Eggleton et al., 1987; Marsh, 1991; Price et al., 1991; Nesbitt and Wilson, 1992; Karrat et al., 1998; Rudnick et al., 2004; Retallack, 2008; Oh and Richter, 2005; Hausrath et al., 2011; Sanematsu et al., 2011; Liu et al., 2013). In Section 5.4, correlations of the weathering indices with the loss on ignition further demonstrate the mineralogical transformations that dictate the chemical weathering trends.

5.1. The chemical index of alteration (CIA)

5.1.1 Quantifying the intensity of weathering with the CIA

Most igneous rocks of varying composition will plot between a CIA value of 35 and 50, with mafic rocks occupying the lower values. Similar to other basalts, the least-weathered (parent rock) samples of both the Chhindwara and Bidar profiles yield a CIA value of ~35.
During chemical weathering, the values of weathering indices increase due to the loss of mobile elements relative to an element assumed to be immobile. The CIA predominantly tracks feldspar dissolution and the concomitant release of Ca, Na, and K relative to Al, since the latter is typically retained within pedogenetic clays. This is evident in the clear separation of the CIA values of primary minerals (e.g., plagioclase, pyroxene: 50 and under) from those of pedogenetic minerals such as smectites, illite (70-85), and kaolinite (100).

In the Chhindwara profile, the samples extend from the unweathered CIA value of 35 to a maximum of 80 in their most altered state. This indicates that the most weathered samples of the Chhindwara profile have not yet reached complete ‘kaolinitisation’ and still retain a detectable amount of the labile elements (Ca, Na, and minor K), possibly due to the presence of zeolite minerals (Bhattacharyya et al., 1999). By contrast, the altered samples of the Bidar laterite profile have CIA values of 90 or greater. These high CIA values indicate the near-complete removal of labile major element cations. The CIA is ineffective at quantifying or differentiating elemental changes during the advanced stages of weathering since the dominant process occurring during lateritisation is desilication and Si is not factored into the CIA.

5.1.2 The A–CN–K diagram

The greatest strength of the CIA as a chemical weathering proxy is the utility of the accompanying A–CN–K diagram, which has empirically and kinetically predictable weathering vectors for various minerals and rock types (e.g., Nesbitt and Young, 1984; Nesbitt, 1992). The overall weathering vector for feldspar destruction in various different parent rocks is parallel or sub-parallel to the A–CN axis (e.g., Figure 5a). The precise vector direction is a function of the relative proportion of plagioclase and K-feldspar, their congruent or incongruent dissolution, and
the rate of conservation of aluminous weathering products. As predicted for basalt weathering (Nesbitt and Wilson, 1992), the majority of the Chhindwara samples follow a weathering trend that is adjacent to the A–CN join (Figure 5a). Should the degree of weathering continue, the vector is predicted to continue along the A–CN join until it reaches the A apex and the Ca, Na, and K is completely removed (complete ‘kaolinitisation’). Further chemical weathering trends cannot be represented in the A–CN–K plot, as demonstrated by the Bidar laterite samples, which cluster around the A apex (Figure 5a).

5.1.3. K enrichment

A further utility of the A–CN–K plot is identifying and unravelling the effects of potassium enrichment in weathering profiles, such as the conversion of smectite group minerals to illite during diagenesis (Fedo et al., 1995). Enrichment of K results in a trend away from the weathering vector towards the K apex. This is evident in three samples located near the contact of the lower flow (ChQA) in the Chhindwara profile (K concentration of 1.7-1.8 wt. %). The position of these samples deviates from the main weathering trend in the A–CN–K plot defined by the remainder of the profile (Figure 5a). The significance of these samples is discussed further in Section 6.2.3.

5.2. The mafic index of alteration (MIA)

5.2.1 Introduction to the MIA

The mafic index of alteration (MIA) is proposed here as a chemical weathering index that extends the equation of the CIA to include the mafic elements Mg and Fe. Many of the mafic minerals (pyroxene, olivine) are susceptible to chemical weathering, resulting in the loss of Mg from weathering profiles. The loss of Mg can be monitored independently using the Mg index
By contrast to Mg, the fate of Fe during the weathering of most mafic minerals is redox-dependent. In reducing environments, ferrous iron $[\text{Fe}^{2+}]$ can be mobile and leached along with Mg during mafic mineral weathering. In oxidative weathering environments, however, Fe is usually retained by the formation of highly insoluble ferric iron $[\text{Fe}^{3+}]$ oxides or oxyhydroxides (e.g., Driese, 2004) and thus enriched along with Al. Due to this dichotomous redox behaviour of Fe, an arrangement of the MIA is proposed for each of the end-member weathering environments.

When the alteration environment is oxidising and Fe is retained, total Fe is considered an immobile element along with Al ($\text{Al}_2\text{O}_3$) and the MIA calculation is:

\begin{equation}
\text{MIA}_{(O)} = 100 \times \left( \frac{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(T)}{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(T) + \text{MgO} + \text{CaO}^* + \text{Na}_2\text{O} + \text{K}_2\text{O}} \right)
\end{equation}

When the alteration environment is reducing and Fe is leached along with Mg, total Fe is considered a mobile element along with Mg, Ca, and K and the MIA calculation is:

\begin{equation}
\text{MIA}_{(R)} = 100 \times \left( \frac{\text{Al}_2\text{O}_3}{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(T) + \text{MgO} + \text{CaO}^* + \text{Na}_2\text{O} + \text{K}_2\text{O}} \right)
\end{equation}

In both arrangements of the MIA [$\text{MIA}_{(O)}$ or $\text{MIA}_{(R)}$], increasing index values represent progressively more altered rock, as is the case with the CIA. A value of 100 indicates complete removal of the mobile elements. As with the CIA, the MIA uses the molar ratios of the major element oxides by converting the wt. % concentrations into moles (see Table 3). The molar CaO is corrected for the presence of carbonate and apatite as for the CIA (e.g., Fedo et al., 1995) to consider only the silicate-bound Ca ($\text{CaO}^*$).

In addition to the index value, the proportional changes of the elements in the MIA can be studied graphically with ternary plots in the Al–Fe–Mg–Ca–Na–K system (e.g., Nesbitt and
Young, 1989; Nesbitt and Wilson, 1992) whereby both versions of the MIA are arranged into the plots in the same manner as the CIA with the A–CN–K diagram. The MIA\(_{(R)}\) arranges into the

\[
\text{Al}_2\text{O}_3–(\text{CaO}^* + \text{K}_2\text{O} + \text{Na}_2\text{O})–(\text{Fe}_2\text{O}_3(T) + \text{MgO}) (A–CNK–FM) \text{ plot (Nesbitt and Young, 1989),}
\]

while the MIA\(_{(O)}\) arranges into the \(\text{Al}_2\text{O}_3–(\text{CaO}^* + \text{MgO} + \text{Na}_2\text{O} + \text{K}_2\text{O})–\text{Fe}_2\text{O}_3(T)\) (A–L–F) plot (Nesbitt and Wilson, 1992) as well as the newly proposed \((\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(T))–(\text{CaO}^* + \text{Na}_2\text{O} + \text{K}_2\text{O})–\text{MgO} (AF–CNK–M) \text{ plot. The details of these plots are presented in Sections 5.2.3–5.2.5.}

It needs to be cautioned that the MIA calculation values and the trends in ternary plots will vary slightly depending on the use of FeO or Fe\(_2\)O\(_3\) for total Fe due to the cation mole difference. The factor for converting between FeO wt. % and Fe\(_2\)O\(_3\) wt. % is shown in Table 3. Either oxide can be applied in the MIA, but consistency should be used between the weathering index calculations and the ternary plots and when comparing separate data sets. In this study, the moles of Fe are calculated using the total Fe as Fe\(_2\)O\(_3\). All literature values used were recast to total Fe expressed as Fe\(_2\)O\(_3\) and all of the major element oxides subsequently normalised to 100% on an anhydrous basis to allow for accurate comparison.

The application of the MIA and associated ternary plots to studying weathering profiles is best accompanied with independent knowledge of the redox-related behaviour of Fe. This can be accomplished with independent tests of iron immobility or the determination of the Fe\(^{2+}\) content. For example, a ternary plot of \(\text{Al}_2\text{O}_3–\text{MgO–Fe}_2\text{O}_3(T)\) (A–M–F) or a cross-plot of the CIA vs. Fe\(_2\)O\(_3(T)\) can be used to examine the net behaviour of iron (i.e., retention or loss) in a weathering profile relative to Al, as demonstrated by Young (2013). Plots of total Fe vs. Ti or Al have also been used to test for iron mobility (e.g., Rye and Holland, 1998).

5.2.2 Comparison of the MIA and CIA
There is an inherent range in MIA and CIA values for unweathered igneous protoliths reflecting bulk composition (see supplementary information). For the purpose of this study, discussion is limited to mafic compositions and their weathered products. The weathering index values of several mafic USGS standards and parent rock samples from the Bidar and Chhindwara profiles are summarised in Table 4. In all cases, the MIA(O) values are low (less than 45) and close to the calculated CIA values. The CIA vs. MIA(O) values from the Bidar and Chhindwara profiles are plotted in Figure 6a along with the compilation of sub-Recent mafic and Tertiary laterite/bauxite profiles. Overall, the MIA(O) is only slightly lower than the CIA for the majority of samples and the data are very well correlated ($r^2=0.900$), confirming the similar bulk weathering behaviour of Mg, Ca, and Na. The overall correlation between the two indices is poorest between a CIA value of 60 and 85, which could indicate a slightly different behaviour of Mg from Ca and Na during pedogenetic clay formation in the intermediate weathering stages.

When examined in greater detail, it is evident that the MIA(O) and CIA are nearly linearly correlated in the upper flow of the Chhindwara profile ($r^2=0.978$; up to a CIA and MIA value of approximately 70), but decoupled in the lower flow. In the latter, the CIA continues to increase to a value of 80 while the MIA(O) remains relatively static between 68 and 70, suggesting a divergence in the behaviour of Mg and/or Fe from Ca and Na. The concentrations of Fe and Ti are correlated and the $Fe^{2+}/Fe^{3+}$ ratio decreases from 2.56 to 0.03 with increasing CIA values (Figure 6b) in the entire profile, suggesting complete Fe retention via oxidative weathering.

Therefore, it appears that the CIA-MIA(O) decoupling is most likely related to minor Mg retention in the samples from which Ca and Na are more heavily depleted, perhaps in a smectite group mineral or zeolite. Consequently, the MIA is less sensitive to the chemical changes that occur in the lower flow of the profile.
In the Bidar profile, the MIA values are high and relatively invariable (from 91 to 99), similar to the CIA, showing that the MIA suffers from the same inability to quantify the advanced to extreme stages of weathering.

5.2.3 The A–CNK–FM diagram

The A–CNK–FM ternary diagram was proposed by Nesbitt and Young (1989) in order to consider the mafic mineral component in rock weathering. In this plot (Figure 7a), the feldspar (felsic) weathering vector should follow a direction approximately away from the CNK apex through the position of the unweathered sample. The exact magnitude and direction is largely influenced by the rate of feldspar weathering and the type of secondary weathering product. The loss of Fe and Mg from the mafic minerals should result in a weathering vector that emanates away from the FM apex through the unweathered sample. During oxidative weathering, however, Fe is not typically lost from the system. This complicates the mafic weathering trends and contributes to the difficulty in predicting the overall rock weathering vectors (Nesbitt and Young, 1989; Nesbitt, 1992). Regardless, empirical data for Phanerozoic basalt weathering (e.g., Nesbitt and Wilson, 1992) show that progressive alteration of basalt results in a trend away from the CNK apex. Such a trend is evident in the Chhindwara weathering profile (Figure 7a). Taken alone, a net weathering vector emanating away from the CNK apex suggests a greater relative mobility and more extensive leaching of Ca, Na, and K compared to Mg. This may be misleading, however, since Mg is known to exhibits a similar bulk weathering behaviour to Ca and Na (Nesbitt and Wilson, 1992). Therefore, it appears that the behaviour of Fe and Mg are not accurately represented in the plot due to their contrasting behaviour in most (oxidative) weathering environments. It is not until Ca, Na, and K are depleted and the stages of advanced weathering are reached that change in the relative proportion of Mg, Fe and Al are recorded on
the A–CNK–FM plot (Nesbitt and Young, 1989). This is demonstrated by the lateritised Bidar samples in Figure 7a, which fall on the A-FM axis and are a function of the relative proportion of aluminous (e.g., gibbsite, kaolinite) and ferruginous (e.g., hematite, goethite) pedogenetic products.

Our analyses suggest that the A–CNK–FM diagram is most applicable to weathering environments in which Fe$^{2+}$ is mobile and behaves like Mg during chemical weathering (i.e., in reduced geochemical environments, such as recorded in Archaean and early Proterozoic weathering profiles). In these environments, loss of Mg and Fe is summative and results in a mafic mineral weathering vector away from FM apex. Ultimately, this would result in a net weathering vector that is directed away from the CNK–FM axis towards the A apex. In this scenario, the MIA$_{(R)}$ can be projected into the diagram and is equivalent to the tie-line between the A–CNK and A–FM axes, as shown in Figure 7a. In oxidative weathering environments, one of the two following ternary re-arrangements is more useful.

5.2.4 The A–L–F diagram

One solution to the competing Fe and Mg vectors of the A–CNK–FM diagram, is the A–L–F diagram of Nesbitt and Wilson (1992), where Mg is moved to the same apex as Ca, Na, and K (Figure 7b). These four elements comprise the total labile (L) cation content. The upper apex is Al and the remaining axis becomes total Fe. In this plot, the net rock weathering vector is predicted to emanate away from the L apex through the position of the unweathered sample as Ca, Na, K, and Mg are lost. The magnitude of this vector represents the degree of element loss during weathering. This trend is demonstrated with the Chhindwara profile and the literature compilation in Figure 7b. Minor differences in the starting modal mineralogy and the exact
pedogenetic weathering products, however, appears to result in slightly different slopes in the empirical weathering vectors. Complete loss of these elements would result in the trend reaching the A–F axis. Accordingly, the MIA(O) value in the A-L-F plot is equivalent to the tie line joining the L–A and L–F axes, with values increasing towards the A–F tie axis (MIA value of 100).

Although MIA(O) values reach a maximum at the A-F axis, one advantage of the A–L–F plot is that advanced weathering trends can be graphically visualised by trends moving towards the Fe apex (lateritisation) or the Al apex (bauxitisation). The proportion of the aluminous and ferruginous minerals in the advanced weathering residue can be extracted from the plot based on the sample position trend along the A-F axis (Nesbitt and Wilson, 1992). For example, the Bidar samples plot along the A-F axis moving towards the F apex, demonstrating the Fe enrichment in the profile (Figure 7b). Therefore, the plot is also useful for assessing the net behaviour of Fe (i.e., enrichment or loss) during weathering in modern and ancient weathering profiles, similar to the A-M-F plot (Young, 2013).

5.2.5 The AF–CNK–M diagram

For further assessing (oxidative) mafic weathering trends, another complementary ternary plot is proposed: the AF–CNK–M ternary plot (Figure 7c). This ternary diagram is generated by a simple rearrangement of Fe to the upper apex along with Al, leaving Mg to its own apex. As such, the plot allows the immobile behaviour of Fe in oxidised weathering environments to be separated from the typically mobile behaviour of Mg. The main advantage of this arrangement of the apices is that the relative contribution of CNK (feldspar) dissolution to the overall weathering vector can be compared to that of Mg loss. The net weathering vector is the summation of the two independent vectors emanating away from the CNK and M apices towards the AF apex. This
is demonstrated with the literature compilation in the plot (Figure 7c). On the scale of an
individual weathering profile, the starting position and exact vector direction is influenced by the
proportion of mafic minerals to feldspar and the rate at which they weather relative to each other.
In the AF–CNK–M plot, the MIA is equivalent to the tie line between the CNK and M apices,
increasing upwards toward the AF apex (MIA value of 100).

The samples from the Chhindwara profile on the AF–CNK–M plot illustrate that plagioclase
weathering (loss of Ca, Na) has exceeded mafic mineral weathering (loss of Mg), resulting in an
empirical trend that is directed slightly towards the AF-M join. However, by comparison with the
A–CNK–FM plot, the AF–CNK–M plot confirms that Mg does contribute significantly to the
weathering vector, since the empirical trend is not directed purely away from the CNK apex.
Furthermore, the AF–CNK–M emphasises the change in weathering behaviour of Mg in the
lower flow of the Chhindwara profile compared to the upper flow (Section 5.2.2); in the lower
flow samples, the loss of Mg appears to slow while minor Ca and Na loss continues as indicated
by the change to a horizontal vector in the diagram (Figure 7c).

In the AF–CNK–M plot, advanced weathering trends converge towards the AF apex and the
enrichment of an Al vs. Fe cannot be diagrammatically visualised as it can in the A–L–F plot.
Therefore, all of the Bidar laterite profile samples plot at or near the AF apex at MIA_{(O)} values
greater than 90.

5.2.6 Extended application and limitations of the MIA

The focus of the present study is to highlight the applicability of the MIA to quantifying
weathering intensity and associated chemical changes in mafic substrates. Apart from this
application, the use of the MIA may extend to studying intermediate and felsic rock weathering,
sediments and sedimentary rocks, or different types of alteration. It needs to be strictly noted, however, that unweathered igneous rocks of different composition will have varying initial MIA(O) and MIA(R) values (Supplementary information). Therefore, a singular MIA value should not be used to infer the extent of weathering unless some knowledge of the parent rock to a weathering profile or sediment is known.

One of the unique applications of the MIA is for quantifying chemical weathering trends in the ancient paleosol record, since the MIA(R) can be applied to paleosols in which Fe is lost (anoxic paleosols), while the MIA(O) can be used for those in which Fe is retained (oxygenated paleosols). Similar to modern weathering environments, most Precambrian paleosols and clastic sedimentary rocks are significantly depleted in Ca and Mg and, thus, the MIA may be useful in the modelling of global chemical weathering fluxes in both the modern and ancient rock record (e.g., Kramers, 2002). Additionally, both versions of the MIA can be calculated without K (MIA-K), in a manner analogous to the CIA, to counter the effects of K metasomatism (e.g., Fedo et al., 1995; Maynard, 1992). Caution must be used when inferring chemical weathering intensity in rocks that have experienced post-weathering Fe and/or Mg mobility.

5.3 The index of lateritisation (IOL)

It is clear that the MIA(O) and CIA are incapable of adequately quantifying the stages of advanced chemical weathering. A different approach is required that is suited to the chemical changes in laterite or bauxite profiles. During lateritisation, dissolution of quartz and kaolinite (congruent or incongruent) and the enrichment of Fe oxides are the dominant processes, resulting primarily in a loss of Si relative to Al and Fe in the weathered residue (e.g., Widdowson and Gunnell, 1999; Hill et al., 2000).
Schellman (1981, 1982, 1986) proposed a chemical classification of laterites based on the SiO$_2$:($\text{Al}_2\text{O}_3+\text{Fe}_2\text{O}_3$) ratio and the SiO$_2$–$\text{Al}_2\text{O}_3$–$\text{Fe}_2\text{O}_3$($\text{T}$) (SAF) ternary plot to distinguish between kaolinitised, lateritised, and bauxitised weathering residues. First, a notional ‘limit of kaolinitisation’ for the early to intermediate stages of weathering can be calculated, which is associated with relatively minor Si loss. Desilication begins to dominate beyond this limit and a quantitative tripartite progression of the ‘degrees’ (i.e., weakly, moderately, and strongly) of lateritisation or bauxitisation can be made. This chemical classification, however, is not unanimously accepted (e.g., Bourman and Ollier, 2002; 2003) primarily because accurate representation of the alteration progression is premised upon the autochthony of a weathering profile (Figure 1) and there is a necessity to identify the protolith in order to complete the calculation. Therefore, there is potential of its inappropriate use without this type of thorough geological contextualisation. However, where applied in combination with careful geomorphological and geological interpretation, this method of chemical classification does offer significant insight into the alteration progression in a weathering profile (Schellman, 2003; Widdowson, 2007). Accordingly, we promote use of the SiO$_2$:($\text{Al}_2\text{O}_3+\text{Fe}_2\text{O}_3$) ratio and here suggest it should be referred to as the ‘index of lateritisation’ (IOL) or the ‘index of bauxitisation’ (IOB). To be consistent with the chemical classification of laterite and the SAF plot, we define the IOL using the mass (wt. %) ratio of SiO$_2$, $\text{Fe}_2\text{O}_3$($\text{T}$), and $\text{Al}_2\text{O}_3$, as shown in equation 4.

\[
\text{Eq. 3: IOL} = 100 \times \left[ \frac{(\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(\text{T}))}{(\text{SiO}_2 + \text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(\text{T}))} \right]
\]

The IOL value is designed to directly accompany the SAF diagram, similar to the approach of Hill et al. (2000). Unweathered mafic rocks have IOL values that are generally less than 40, as demonstrated by the mafic USGS standards and the least-weathered samples in this study (Table...
for instance, unweathered Deccan basalt (BB1) gives an IOL value of 35.7. In a similar fashion to the CIA and MIA, higher IOL values correspond to more intensely weathered samples; accordingly, the calculated IOL values for increasingly weathered samples progressively fall within the divisions of kaolinitisation, weak, moderate, and strong lateritisation, as determined from the composition of the protolith. The aforementioned divisions are calculated based on the amount of Si necessary to convert Al in the protolith into kaolin (see Widdowson, 2007, and the method given in the supplementary materials). The limit of kaolinitisation for the Deccan basalt, calculated from sample BB1, is 43% SiO$_2$ (Figure 5). In terms of the IOL, this division between kaolinitisation and lateritisation occurs at an IOL value of 57; for reference, both BB3 and BB4 lie near this boundary (i.e., IOL value of 59 and 58, respectively).

In the Chhindwara profile, samples are restricted to IOL values of ~35-50, consistent with the modest loss of SiO$_2$ relative to Al$_2$O$_3$ and Fe$_2$O$_3$ during kaolinitisation. The Bidar laterite, however, extends across the kaolinitisation-lateritisation boundary and defines a clear trend towards the F apex with IOL values ranging from 57-94 (Figure 5b; Borger and Widdowson, 2001, Widdowson, 2007). The more extreme range in the IOL values is consistent with the upward increase in modal abundance of Fe oxides in the Bidar profile. As discussed previously, however, the samples BB5 and BB6 do not conform to the upward Fe-enrichment trend held by the remaining profile. These samples (~10-15 m depth; Figure 3), interpreted to represent a paleo-water table (e.g., Kisakürek et al., 2004; Widdowson, 2007), contain significantly higher Fe (and other metal) concentrations than would be expected at their stratigraphic position. Therefore, comparing the stratigraphy with the IOL may help to identify samples that do not meet the criteria for strict *in situ* formation.
5.4 Loss on ignition

The volatile content of a sample measured by the loss on ignition is considered proportional to the amount of hydrated minerals, in addition to carbonate, organic carbon, sulphur, less any mass gain from oxidation of elements during analysis. The LOI value of a fresh basalt is typically less than 3 wt. %. During chemical weathering, as the primary and largely anhydrous magmatic minerals are replaced by hydrated pedogenetic phyllosilicates (e.g., smectites, illite, kaolinite), the LOI is expected to increase. Previous studies have shown that the LOI can be a useful parameter for assessing the degree of weathering (e.g., Duzgoren-Aydin and Aydin, 2003), although the specific pedogenetic mineralogy ultimately controls the degree of hydration (Duzgoren-Aydin et al., 2002). Plotting the CIA, MIA(O), or IOL weathering indices of the Chhindwara and Bidar profiles against the LOI provides further insight into the pedogenetic mineralogical transformations (Figure 8).

A distinct positive correlation between the LOI (~1 to 18 wt. %) and the MIA(O) or CIA weathering indices is exhibited by the Chhindwara samples (Figure 8a). By contrast, the more intensely weathered samples of the Bidar profile are removed from the correlation and are less hydrated, with LOI values ranging from ~7 to 12 wt.%. Whereas this may seem counter-intuitive, the lateritised samples are dominated by gibbsite, goethite, and hematite (Kisakürek et al., 2004) which are less hydrous than the intermediate weathering products (kaolinite, smectites). The replacement of the hydrated phyllosilicates by sesquioxides of Fe and Al during lateritisation results in an inverse correlation with the LOI (Figure 8b). In effect, this observation provides mineralogical corroboration of the SAF plot since the change in weathering vector in Figure 5b effectively represents the notional ‘limit of kaolinitisation’ determined for the parent basalt.
6. RESULTS AND DISCUSSION – PART II

Mass gains and losses of the alkali and alkaline earth elements

The following sections discuss the geochemical behaviour of the alkaline earth (Section 6.1) and alkali elements (Section 6.2) during basalt weathering. The analysis employs a combination of mass balance values, element-element ratios, and the major element chemical weathering indices. Combining the independently calculated mass changes with the chemical weathering indices has the power of linking high-precision trace element data with specific stages of basalt alteration. The discussion will only consider the Chhindwara weathering profile since the elemental systematics of the Bidar laterite have previously been analysed (e.g., Kisakürek et al. 2004) and have been complicated by ancient paleo-water table changes and aeolian input (Mason et al., 2000; Kisakürek et al., 2004; Wimpenny et al., 2007). The K-enriched samples in the lower flow of the Chhindwara profile (Section 5.1.3) are discussed separately from the remaining samples in Section 6.2.3. The release or retention of elements in the profile is linked when possible with the Deccan Traps river water (Das et al., 2005) and sediment (Das and Krishnaswami, 2006, 2007) chemistry that is sourced from the DVP basaltic terrane.

For mass balance calculations, the tau (τ) mass transport model (Brimhall and Dietrich 1987; Anderson et al., 2002) is used. In this model, the concentrations (C) of elements (j) in the parent rock (p), relative to that of an immobile index element (i), are used as a normalisation to establish the mass changes in the progressively altered rock (w):

\[ \tau_{ij} = \frac{[(C_{j,w}) / (C_{j,p})]}{[(C_{i,w}) / (C_{i,p})]} - 1 \]
The sample representing the parent rock of the Chhindwara weathering profile is ChQB12 (depth of 140 cm) based on its low degree of chemical alteration (CIA: 36, MIA\(_\text{O}_{\text{O}}\): 38) and Nb is used the immobile index element (Widdowson and Cox, 1996). The \(\tau_{\text{Nb},j}\) values are plotted vs. depth in Figure 9. It should here be remembered that the Chhindwara profile is developed across two independent lava flows. Each flow can be fingerprinted chemically using immobile element ratios such as Al\(_2\)O\(_3\)/TiO\(_2\); the upper flow has a mean ratio of 5.74 ± 0.35 whereby the lower flow is distinct with a mean ratio of 4.27 ± 0.19. Normalisation using the immobile element composition of the parent material in the upper flow results in minor mass balance offsets (\(\tau_{\text{Nb},j}\) values) for elements in the lower flow that are unrelated to pedogenesis. This does not significantly influence the interpretation of the mass balance changes, but a slight inaccuracy in the reported mass gains and losses is expected as a result.

### 6.1 Alkaline earth elements (+ Na)

The elements of the alkaline earth group are the most mobile during continental weathering (e.g., Nesbitt et al., 1980), being hosted in mineral phases most susceptible to chemical attack and, in general, incompatible in pedogenetic clays. In Figure 10, the \(\tau_{\text{Nb},j}\) values of the alkaline earth elements are plotted against the CIA and an anti-correlation exists for nearly all of the elements. The degree of element depletion (represented by the steepness of the slope) differs as a function of how closely related the behaviour of the element in question is to Na and Ca.

Within a ‘stratigraphic’ context, the degree of alkaline earth element (+ Na) depletion in the Chhindwara flows follows the visible extent of basalt alteration (Section 3.3); the \(\tau_{\text{Nb},j}\) values for the alkali elements change only minimally in the centre of the upper flow, but decrease near the modern soil surface and at the flow-banded area at the base (Figure 9). Within the lower flow,
the degree of depletion and CIA values are higher and increase from the deepest sample 
(ChQA1) towards the flow contact (ChQA10-12) in parallel with the vesicle (amygdale) 
abundance. The overall weathering behaviour of the alkaline earth elements (+ Na), assessed 
using the τ mass transport model and molar element ratios, suggests an order of susceptibility of 
Na ≈ Ca ≈ Sr > Mg > Ba > Be. This order of mobility during weathering, with the exception of 
Mg, is consistent with the data from work of on the river sediments and water of the drainage 
basins in the Deccan Traps (Das et al., 2005; Das and Krishnaswami, 2006, 2007), as discussed 
further below.

6.1.1 Na, Ca, Sr

The behaviour of Ca, Na, and Sr are strongly linked in the profile. For example, the Sr 
concentration is highly correlated with the CIA ($r^2=0.977$) and the $\tau_{Nb,Sr}$ values with $\tau_{Nb,Ca}$ 
($r^2=0.973$). These elements exhibit the greatest net depletion based on the $\tau_{Nb,j}$ values (Na: 100%, 
assuming a concentration of 0 for analyses below the detection limit of the XRF; Ca: 90%; Sr: 
87%). They are hosted predominantly in plagioclase, which is highly susceptible to chemical 
weathering. The average molar Ca/Na ratio of the least-weathered basalt samples (CIA ≤ 40: 
ChQB12, ChQB9b-d) in the Chhindwara profile is 2.26 ± 0.06 (n=4), which is near the average 
of the Poladpur and Ambenali formations (2.45-2.53; Subbarao et al., 2000; Widdowson et al., 
2000). The Ca/Na is relatively constant in the remainder of the profile, with two exceptions. 
First, slightly lower Ca/Na ratios are present in three samples at the base of the upper flow 
(ChQB3-ChQB5; Ca/Na: 1.60-1.75), possibly indicating enhanced Ca removal. Second, the 
Ca/Na ratios approach infinity for the samples in which Na is below the detection limit of the 
XRF. This observation, coupled with the near constant molar Ca/Na ratios in Deccan river
sediments (Das and Krishnaswami, 2007) and river waters (Das et al., 2005) suggests that Ca and Na are removed congruently from the Deccan basalt during chemical weathering.

6.1.2 Mg

A strong depletion of Mg follows the other alkaline earth elements at the base of the upper flow, as indicated by the correlation of $\tau_{\text{Nb,Mg}}$ with $\tau_{\text{Nb,Ca}}$ ($r^2=0.952$) and the relatively invariable molar Ca/Mg ratio. The latter ratio is similar to the least-weathered basalt samples (constant at $1.37 \pm 0.03$, $n=4$) and the range exhibited by unweathered Deccan basalts (Subbarao et al., 2000; Widdowson et al., 2000). The change in weathering behaviour of Mg in the lower flow (Section 5.2.2) is expressed by lower Ca/Mg ratios, which range from 0.29 to 0.72. Although the spread in $\tau_{\text{Nb,Mg}}$ values in the lower flow is small (65-73% depletion), the minor variations appear to be anti-correlated with Ca, Na, and Sr loss. This change in Mg weathering behaviour is unlikely to be important to the overall flux of Mg to the hydrosphere from the weathering of the Deccan basalt province. Near-constant Mg/Sr and Ca/Mg ratios of the Deccan river sediments suggest a similar and congruent weathering behaviour of Mg to these other alkaline earth elements (Das and Krishnaswami, 2007). However, the river water data do exhibit notable scatter in the Ca/Mg ratio (Das et al., 2005) and this may indicate that preferential leaching of Ca relative to Mg is important locally, consistent with the observations of Mg in the lower flow of the Chhindwara profile.

6.1.3 Ba

The $\tau_{\text{Nb,Ba}}$ values, although reaching up to 68% depletion, indicate a higher retention for Ba than most other alkaline earth elements and the values are less correlated with the CIA. There are some areas of local Ba enrichment (~75%), such as near the soil surface. By comparison, the
ionic radius of Ba$^{2+}$ is larger and much closer to that of K$^+$, suggesting a stronger association with pedogenetic phyllosilicates. Although Ba and K are not directly correlated in the Chhindwara profile, they are in the river sediments of the Deccan Traps (Das and Krishnaswami, 2007), indicating a broadly similar retention in sediments derived from the Deccan Traps. Barium has long been known to exhibit a different weathering behaviour than the other, more mobile alkaline earth elements (e.g., Nesbitt et al., 1980; Buggle et al., 2011). It is often contrasted with Sr since the Ba$^{2+}$ and Sr$^{2+}$ ionic radii differ by roughly 12-13% in the same coordination, leading to the disparity in their mobility. Accordingly, the Ba/Sr ratio has been applied as an index of the degree of alkaline earth element depletion or leaching during hydrolysis (e.g., Retallack, 1994; Gallet et al., 1996). The Ambenali and Poladpur formations have average molar Ba/Sr ratios that range from 0.24 to 0.38 (Widdowson et al., 2000). The Ba/Sr ratio of the Chhindwara profile evolves from a similar ratio of ~0.3 in the least-weathered samples to a maximum of 0.7 (if the horizons of Ba enrichment are excluded). Collectively, the Ba geochemistry of Deccan Traps river waters and sediments confirm its less mobile behaviour during basalt weathering. The mean ratios of Ba to other alkaline earth elements (Sr, Mg, Ca) in Deccan rivers are typically 2-3 times lower than the Deccan basalt (Das and Krishnaswami, 2006) and the river sediment Ba/Sr ratios are highly variable, ranging from 0.3 to 3.2, with a mean value of 1.1 (Das and Krishnaswami, 2007). The sediment samples with the highest Ba/Sr appear to be heavily influenced by the sediments of the Krishna tributaries, which may have a greater input of lateritised products that experienced greater degrees of leaching.

6.1.4 Be
In the Chhindwara weathering profile, the $\tau_{\text{Nb,Be}}$ values are the highest of all of the alkaline earth elements, suggesting it is the least mobile of the group. The best correlation of $\tau_{\text{Nb,Be}}$ is found with $\tau_{\text{Nb,Ba}}$ and $\tau_{\text{Nb,Li}}$. Despite its slight depletion, it appears that the small ionic radius of $\text{Be}^{2+}$ results in a less mobile behaviour, possibly due to the greater potential of the cation to form insoluble hydrolysates.

**6.2 Alkali elements (+Thallium)**

The alkali elements, with their lower ionic potential compared to the alkaline earths, are more strongly associated with the phyllosilicates, substituting into interlayer sites or adsorbing to clay mineral surfaces. Despite the high solubility of many of the alkali elements, this factor tends to lead to a lower mobility during weathering (Nesbitt et al., 1980). This is evident by the $\tau_{\text{Nb,j}}$ values for the alkali elements (see Figure 9 for values and trends with depth), which exhibit minimal to no positive or negative correlation with the CIA, indicating a strong decoupling from the mobile alkaline earth elements. When the $\tau_{\text{Nb,j}}$ values of several alkali elements are plotted against each other, two separate correlations normally result, suggesting the mass balance calculations are more greatly influenced by the different parent flow chemistry. Included with the alkali elements is thallium (Tl) due to its similar geochemical properties, namely a commonly monovalent charge and similar ionic radius (Shannon, 1976). The behaviour of Li is largely decoupled from the remaining alkali elements and is discussed separately in Section 6.2.1.

Overall, the alkali elements closest in ionic radius exhibit the most similar behaviour and the correlations between the element concentrations and mass balance values decrease. For example, strong correlations exist between $\tau_{\text{Nb,K}}$ and $\tau_{\text{Nb,Rb}}$ within the separate flows in the Chhindwara profile ($r^2=0.832$ in the upper and $r^2=0.939$ in the lower). Using similar relative relationships, the
behaviour of Cs is most closely associated with Rb, and the behaviour of Tl is closely related with Rb. In the latter case, a similar weathering behaviour can also be inferred from the strong correlation of these elements in alluvial sediment (Kamber et al., 2005).

6.2.1 Li

A more significant anti-correlation, similar to the alkaline earth elements, is found between the $\tau_{\text{Nb, Li}}$ values and the CIA (Figure 10). As an expected consequence, the $\tau_{\text{Nb, Li}}$ values correlate better with those of Na, Ca, and Sr and less with the other alkali elements (i.e., Rb, K, Cs). The behaviour of lithium during continental weathering is reasonably well understood and the subject of more detailed focus due to stable isotope fractionation during pedogenesis (Huh et al., 2001; Rudnick et al., 2004). In general, Li is expected to be more mobile during weathering than the other alkali elements (excluding Na), but has a stronger affinity for phyllosilicates, likely as a result of coupled substitution with $\text{Mg}^{2+}$ for $\text{Al}^{3+}$ in the octahedral sites (Ronov et al., 1970; Anderson et al., 1989; Huh et al., 2004).

6.2.2 Alkali element enrichment

Withheld from the previous discussion were three samples from the lower flow of the Chhindwara profile that appear to have experienced K addition (Section 5.1.3). Based on the $\tau_{\text{Nb,j}}$ values, this K enrichment is up to 400 % of the parent rock concentration. This enrichment extends to the other elements in the alkali group (Figure 9), with the most extreme enrichment exhibited by Cs (~1500 %), followed by Rb (~1000 %) and Tl (~110 %). In many Phanerozoic to modern weathering profiles, K enrichment is attributed to preferential uptake in plants and/or the addition of an allochthonous, K-rich material such as aeolian dust (Sheldon, 2003).
Stratigraphically, these samples constitute the vesicular and brecciated flow top. Accordingly, input of allochthonous material (dust or sediment) to the lava flow during or after emplacement is a likely explanation for the enrichment (e.g., Ghosh et al., 2006). Since the alkali elements are depleted in the mantle source of most basalt and highly enriched in the continental crust, only minor degrees of allochthonous addition could generate significant chemical enrichments.

Further work, such as high-precision Nd isotope analysis could substantiate this hypothesis (Mason et al., 2000). The alternative explanation, biological enrichment of K (e.g., Sheldon, 2003), remains less favourable since the enrichment extends to the other alkali elements and it is localised to only the flow top zone.

7. RESULTS AND DISCUSSION – PART III

Behaviour of the rare earth elements during basalt weathering

The CI-chondrite normalised REE patterns of the three least-weathered samples from the Bidar (BB1) and Chhindwara (ChQB12 and ChQB9d) weathering profiles (Table 4) are nearly identical, differing only in absolute abundance. This justifies their use as parent rock values for normalising the progressively more weathered samples (Figure 11). This normalisation most readily exposes REE mobility (e.g., Laveuf et al., 2008). The REE are known to be variably mobile during weathering and fractionate during pedogenesis (e.g., Nesbitt, 1979; Duddy, 1980; Braun et al., 1993; Cotten et al., 1995; Laveuf and Cornu, 2009 and the references therein; Sanematsu et al., 2011), but there is still little consensus regarding the details of the behaviour.

7.1 Overall REE behaviour
Fractionation of the REE in the Chhindwara and Bidar weathering profiles is evident in the change in abundance and slope of the normalised REE patterns in Figure 11. In most of the Chhindwara samples, the REE are enriched relative to the least-weathered sample (ChQB12) and that retention follows the general order of LREE>MREE>HREE during the weathering progression. This observation is consistent with previous studies regarding the mobility of the REE during pedogenesis (e.g., Laveuf and Cornu, 2009 and the references therein) whereby the LREE are less readily fluid-complexed than the HREE and become enriched in weathering products such as the phyllosilicates (e.g., Nesbitt, 1979; Mongelli, 1993).

The REE abundance of the samples in the Bidar laterite profile changes more significantly relative to the parent rock (Figure 11c). A more pronounced depletion is evident within the highly lateritised (upper 10 m) zone of the profile. If the REE are predominantly associated with clays in the early-forming weathering residue, the advanced stages of Si loss during lateritisation appears to release them from the profile. A higher retention of the LREE, however, is still present in the lateritised samples (Figure 11c). The MREE are more depleted than the LREE and HREE during the advanced stages of weathering. As an exception to the above samples, BB3 has a highly anomalous REE pattern and abundance. It is significantly enriched relative to the parent basalt and has an extremely high HREE/LREE slope. This sample may represent a horizon in which REE (and preferentially the HREE) leached from higher in the profile and accumulated at a site of contrasting pH and/or decreased permeability (e.g., Braun et al., 1998; Patino et al., 2003; Viers and Wasserburg, 2004; Kamgang Kabeyene Beyala et al., 2009), such as near the saprolite-protolith boundary. When studying deep, highly advanced weathering profiles, however, the addition of allochthonous material (e.g., dust) can also potentially influence the chemistry of the profile during long surface exposure times (e.g., Kisakürek et al., 2004;
Wimpenny et al., 2007). Further, constraining the influence of dust addition to the REE chemistry of the Bidar laterite is also difficult without knowledge of the chronology of lava stacking and dust composition at the time of accumulation.

7.2 Sm/Nd fractionation

In addition to the obvious changes in the LREE/HREE abundance during pedogenesis, fractionations between some of the closely spaced REE are also detectable, as exemplified by the Sm/Nd ratio. If both flows are considered together, a decrease in Sm/Nd correlates with increasing weathering intensity ($r^2=0.813$; Figure 12). The parent basalt Sm/Nd ratio of the upper and lower flows appears to have differed slightly, which complicates the superimposed effects resulting from chemical weathering.

These observations suggest that Sm/Nd and, consequently, Nd isotope systematics of altered basalt and derived sediments are sensitive to pedogenetic fractionation. This would not alter the Nd isotope composition of modern profiles or sediments, but fractionation in Sm/Nd would manifest as variations in the Nd isotope composition in ancient weathering profiles. In this sense, the more weathered material would be less radiogenic as a result of the lower Sm/Nd generated during LREE enrichment. This has been exploited to crudely date the age of paleosols since an isochron develops that is proportional to the age of pedogenesis (e.g., Stafford, 2007; Frei and Polat, 2013). Using the Chhindwara profile as an example, the extent of Sm/Nd fractionation in the upper flow would translate to a one $\varepsilon_{Nd}$ unit difference over a billion years. Note that this calculation assumes that all of the change in the $^{147}\text{Sm}/^{144}\text{Nd}$ is generated from pedogenesis (highest ratio in the parent basalt and the lowest in the most weathered material), the initial
parent rock $^{143}\text{Nd}/^{144}\text{Nd}$ is constant, and that the isotope system remains closed after pedogenetic fractionation.

These findings contradict the conclusions of Nesbitt and Markovics (1997), where changes in Sm/Nd were undetectable in weathered granodiorite. Therefore, more extreme LREE fractionation may occur during basaltic weathering by comparison to more felsic and/or coarser grained rocks. At present, it is difficult to attribute this to a property of the parent rock (e.g., grain size, bulk composition, mineralogy), differences in the pedogenetic mineralogy, or the weathering environment (e.g., drainage, organic matter). Regardless, given the high erosion rates of basaltic terrain and their greater abundance on the early Earth, this is an important consideration for global weathering fluxes. Further, it seems that the Sm/Nd (or La/Ce) ratio may not always be an ideal provenance indicator, as advocated, for instance, by Sheldon and Tabor (2009). Tests of immobility should be made, if possible, prior to provenance interpretation.

7.3 Europium anomaly ($\text{Eu}/\text{Eu}^*$)

Europium is the only lanthanide that commonly occurs in a divalent oxidation state and whose behaviour is strongly influenced by plagioclase. This results in the potential for Eu to fractionate from the other lanthanides during weathering, since plagioclase is one of the most susceptible minerals to chemical dissolution. Fractionation of Eu can be tracked using the Eu anomaly $[\text{Eu}/\text{Eu}^* = \text{Eu}_n/(\text{Sm}_n \times \text{Gd}_n)^{1/2}]$. In the following discussion, the Eu/Eu* values are calculated using normalisation to the parent rock sample of the profile in question. Qualitatively, it is apparent that nearly all of the samples within both weathering profiles have a negative Eu anomaly (Figure 11).
The Eu/Eu* values in the Chhindwara profile decrease from 1 to ~0.75. Importantly, this variation in the Eu/Eu* value is inversely correlated with the CIA (Figure 13a). During the earliest stages of weathering recorded within the upper flow of the Chhindwara profile (CIA: 35-70), there is only a subtle change in the Eu anomaly (to as low as 0.90). Within the lower flow of the Chhindwara profile (CIA: 70-80), the slope of the CIA vs. Eu/Eu* anti-correlation steepens and Eu/Eu* reaches its lowest values. The same inverse correlation of the Eu/Eu* value with the CIA in the Chhindwara profile also exists with Sr concentration and the Rb/Sr ratio (Figure 13b), including the same inflection point at the transition from the upper to lower flow. Although these observations could indicate more aggressive Eu loss at higher degrees of weathering intensity in the lower flow, the minor variation in the primary chemistry of the two flows is probably the cause for the apparently different slope. Regardless, these observations confirm a strong relationship between Eu loss and plagioclase weathering. Similar reports of a declining negative Eu anomaly as a function of weathering intensity have been made previously (e.g., Condie et al., 1995; Huang and Gong, 2001; Ma et al., 2011). In the study of Ma et al. (2011), pore waters within the local weathering residuum possessed a positive Eu anomaly. Lawrence et al. (2006) noted that when river waters were normalised to the composition of their catchment geology, significant variation in the Eu anomaly remained that suggested a more complicated and possibly mineral-specific weathering contribution to the waters (such as preferential plagioclase dissolution).

In the Bidar laterite samples, the Eu/Eu* values are essentially constant between 0.85 and 0.90, indicating that a similar, albeit less pronounced, depletion of Eu has occurred relative to the parent sample (BB1). The Eu/Eu* value, however, does not change as a function of the weathering intensity in the Bidar samples (i.e., there is no obvious correlation between the IOL
and Eu/Eu*). This confirms that the major loss of Eu occurs during the earlier stages of weathering. By the time all plagioclase has been removed from the weathering profile, the Eu/Eu* variability ceases and the remaining Eu changes only as a function of the processes affecting the remaining trivalent lanthanides. The possibility needs to be considered that Eu/Eu* could also have been affected by dust introduction during weathering exposure.

7.4 Ce anomaly (Ce/Ce*)

Cerium can track redox-related transformations during pedogenesis in modern and ancient weathering profiles as a result of the potential oxidation of Ce$^{3+}$ to Ce$^{4+}$ (e.g., Middelburg et al., 1988; Braun et al., 1990; Mongelli, 1993; Gallet et al., 1996; Murakami et al., 2001; Patino et al., 2003). A positive Ce anomaly [Ce/Ce* = Ce$_n$/(La$_n$ x Pr$_n$)$^{1/2}$] develops at sites of Ce enrichment, although there are often cases of weathered samples displaying distinct negative Ce anomalies, presumably generated by the cycling of REE away from areas of Ce enrichment (e.g., Nesbitt, 1979; Marsh, 1991; Fodor et al., 1992; Cotten et al., 1995). It is evident from the REY plots of the weathered Deccan basalt (Figure 11) that Ce is fractionated to varying degrees (relative to La and Pr) in nearly all samples, confirming its decoupling from the other LREE.

The Ce concentration increases during the incipient to intermediate weathering of the Chhindwara basalt along with the other LREE. There is, however, no obvious correlation between the Ce/Ce* value and the degree of weathering and several samples in the upper flow have negative anomalies. The presence of negative and positive Ce anomalies indicates that Ce oxidation is probably occurring at a smaller scale in the weathering front (e.g., Taunton et al., 2000). By contrast, within the lower and more highly weathered flow (CIA > 70), more samples have positive and higher Ce anomalies.
The Bidar laterite profile has much more extreme variation in Ce/Ce* with values ranging from 0.27 to 6.71. The Ce anomaly does not have any obvious correlations with the IOL, although the two highest Ce/Ce* values (BB7: 4.14 and BB9: 6.71) are within the upper, most highly lateritised portion of the weathering profile. This is consistent with the implied importance of specific oxides and secondary minerals (e.g., florencite; Sanematsu et al., 2011) in the fractionation of the REE. The REE-enriched sample deep in the profile (BB3), has the most negative Ce anomaly (0.27), indicating that the lanthanides transported from above were likely leached in the presence of Mn oxides from higher in the profile.

8. CONCLUSIONS

In this study, two weathering profiles located in the Deccan Traps, India, were studied to better quantify and understand different stages of basalt alteration. In basaltic weathering environments, lava flow morphology and texture (e.g., flow banding and vesicularity) are dominant controls on the penetration of weathering fluids and fluid-rock interaction. Mineralogical differences, such as the presence of zeolites, may also affect chemical weathering progression. Weathering profiles can develop across several basalt flows and some primary chemical variations from the original flows may survive overprinting by pedogenesis (e.g., HFSE ratios). From a geochemical perspective, this influences the precision of alteration mass balance calculations, especially where unweathered parent material is not available from different flows.

In order to better understand mafic substrate alteration, a new chemical weathering index, the mafic index of alteration (MIA), is proposed. The dichotomous, redox-related behaviour of Fe is
factored into two separate MIA equations suitable for reducing \([\text{MIA}_\text{R}]\) or oxidising \([\text{MIA}_\text{O}]\) environments. The quantitative MIA value can be used in combination with ternary plots in the Al–Fe–Mg–Ca–Na–K system that display the proportional chemical changes. Like the majority of currently established weathering indices, the MIA is most suitable for understanding the early to intermediate stages of chemical weathering (‘kaolinitisation’). Obtaining a quantitative handle on advanced chemical weathering requires a different approach. We propose using the index of lateritisation (IOL) as a quantitative expression of the Si–Al–Fe (SAF) ternary plot (Schellmann, 1981, 1982, 1986). Plotting chemical weathering indices against LOI values can provide additional insight into pedogenetic mineral transformations.

Chemical weathering studies should always start with the major element composition of the weathering profile prior to trace element or isotopic analysis. When used in combination, weathering indices and high-precision trace element analysis can provide insight into which stages of alteration the trace element loss or retention are associated with. In the present study, the focus is on the alkali, alkaline earth, and rare earth elements of the basaltic weathering profiles. The main observations and conclusions presented are:

- The alkali and alkaline earth element geochemistry in the Chhindwara profile conforms predominantly to the expected weathering behaviour. The alkaline earth elements (+Na) exhibit an order of depletion of \(\text{Na} \approx \text{Ca} \approx \text{Sr} > \text{Mg} > \text{Ba} > \text{Be}\). By contrast, the alkali elements exhibit a greater retention in the altered substrate. Notably, Tl appears to behave most closely to Rb and Li exhibits the greatest mobility of the alkali elements.
- Alkali element enrichment (Cs, Tl, Rb, K) at the paleo-flow top of the Chhindwara profile is best explained by the allochthonous addition of dust to the profile. The results show that the enrichment in K reported in other studies may similarly be accompanied by
the other alkali elements. This may help constrain the mechanism of enrichment (e.g., metasomatism, plant uptake, dust).

- Rare earth element fractionation occurs during weathering of the Deccan basalt whereby the enrichment of the LREE is greater than the MREE and HREE in the weathering residue. Significant pedogenetic fractionation of the closely spaced LREE is also evident from variation in the Sm/Nd ratio. Consequently, this indicates that REE ratios should be treated with caution when used for assessing the provenance of weathered materials.

Fractionation of Sm/Nd can manifest as a varying Nd isotope composition over time.

- The preferential loss of Eu, measured with the Eu anomaly (Eu/Eu*) is highly correlated with Sr, Ca, and Na loss from plagioclase. Following the complete weathering of plagioclase, the Eu/Eu* does not appear to change further as a function of weathering intensity.

- Cerium is fractionated from the trivalent lanthanides during pedogenesis but there is a lack of correlation of Ce/Ce* with any of the weathering indices, consistent with fractionation being controlled by pedogenetic minerals that are not contributing to the calculation of the weathering indices.

The results presented here are relevant to ancient paleosol research and may be useful for studying Martian substrate alteration.

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Brimhall, G.H., Dietrich, W.E., 1987. Constitutive mass balance relations between chemical composition, volume, density, porosity, and strain in metasomatic hydrochemical


**All colour figures are for the web version only**

**Figure 1:** Illustration showing the transitional relationship from end-member autochthonous to allochthonous duricrusts (i.e., laterite to ferricrete), modified from Widdowson (2007). The Bidar (BB) laterite profile is an autochthonous profile formed from *in situ* weathering.

**Figure 2:** Simplified geological map showing the extent of the Deccan Volcanic Province (DVP) within peninsular India and the geomorphology of the weathered basaltic terrane. Inset maps show (a) the mesa topography near Chhindwara in the Madya Pradesh district and (b) the more detailed geology of the SE lobe of the DVP, including the extent of the thick laterite that hosts the Bidar profile. DVP geology maps modified from Borger and Widdowson (2001) and Kisakürek et al. (2004).

**Figure 3:** Geological sketch section of the: a) Bidar laterite (modified from Kisakürek et al., 2004 and Widdowson, 2007) and b) Chhindwara weathering profile. Highlights of the key features in the weathering progression accompany the schematic of each profile. Note the different scales of depth. The Chhindwara weathering profile is developed across two identifiable basalt flows (A – lower flow and B – upper flow). The sample identification and locations are also provided.

**Figure 4:** Field photographs of the Chhindwara weathering profile. (a) The entirety of the sampled weathering profile exposed in a road side quarry. Note the colour change marking the transition from the upper flow (ChQB) to the lower flow (ChQA). (b) Corestone within the upper flow at a depth of 90 cm from the surface that was divided into four sub-samples (ChQBa-d) for geochemical analysis.
Figure 5: (a) Molar Al$_2$O$_3$–CaO$^*$$\text{+Na}_2$O–K$_2$O (A–CN–K) and (b) mass SiO$_2$–Al$_2$O$_3$–Fe$_2$O$_3$(T) (SAF) ternary plots illustrating the different degrees of alteration experienced between the Chhindwara (solid squares) and Bidar (solid circles) weathering profiles. The figure symbols for the Bidar and Chhindwara samples and the data compilation are consistent throughout all subsequent figures unless specified otherwise. In both figures, a data compilation of sub-Recent mafic weathering profiles and Tertiary basaltic laterite and bauxite profiles is plotted as shaded circles (see text for references). The chemical index of alteration (CIA) and the index of lateriti$^*$sation (IOL) may be integrated into both the A–CN–K and SAF ternary plots, respectively, as shown. In the SAF plot, the kaolinitisation and lateritisation boundaries are calculated from the chemistry of BB1 (parent rock of the Bidar profile).

Figure 6: (a) Comparison of the MIA$(_R)$ and CIA weathering indices for the Chhindwara and Bidar profiles, as well as the data compilation from Figure 5. (b) Plot of Fe$^{2+}$/Fe$^{3+}$ vs. CIA for the Chhindwara profile demonstrating the progressive conversion of Fe$^{2+}$ to Fe$^{3+}$ in the basalt with increasing weathering intensity.

Figure 7: Molar ternary plots in Al–Fe–Mg–Ca–Na–K (AFMCNK) space. See Table 3 for molar oxide conversions. The (a) A–CNK–FM and (b) A–CNKM–F (A–L–F) plots were proposed by Nesbitt and Young (1989) and Nesbitt and Wilson (1992), respectively, while the (c) AF–CNK–M plot (shaded) is proposed in the present study. These diagrams expose the contribution of the mafic and felsic mineral components to rock weathering. The integration of the new mafic index of alteration (MIA) into AFMCNK space is illustrated; the A–CNK–FM plot, most suitable for studying weathering in reducing environments, is integrated with the MIA$(_R)$, while the A–L–F and AF–CNK–M plots are integrated with the MIA$(_O)$ for studying oxidative weathering trends.
In addition to the Chhindwara and Bidar samples, the data compilation from Figure 5 is plotted as well.

**Figure 8:** Co-variation of the loss on ignition (wt. %) with the (a) chemical index of alteration and (b) index of lateritisation. The calculated kaolinitisation and lateritisation boundaries from Figure 5b are shown on the latter. Strong correlations of the weathering indices with the loss on ignition during incipient to intermediate weathering (Chhindwara weathering profile) confirm the transformation of relatively anhydrous pyroxene and plagioclase to hydrous phyllosilicates (i.e., smectites and kaolinite). Over the much longer time scales of advanced weathering, Si is lost as these phyllosilicates are progressively altered to sesquioxides of Al and Fe. This trend is defined by a decreasing loss on ignition as the index of lateritisation increases.

**Figure 9:** Stratigraphic variation of the weathering indices (MIA_{10} and CIA) (a) and mass balance (τ mass transport function) profiles of the alkaline earth (b) and alkali elements (c) for the Chhindwara weathering profile. The depth of the modern soil surface and the two individual lava flows are indicated. Mass balance calculations use Nb as the immobile index element and sample ChQB12 (depth of 140 cm) for the parent rock. Values less than 0 indicate a mass loss relative to the parent rock (up to 100 % or -1) while values greater than 0 are mass gains. Note the extreme enrichment in the alkali elements at the top of the lower flow (darker shading).

**Figure 10:** τ_{Nb,j} values vs. the CIA for the alkaline earth elements (plus Na and Li). Note that values below 0 (dotted line) represent a mass loss up to -1 (100 % depletion). The τ_{Nb,j} value of most elements is anti-correlated with the CIA indicating that they are mobile during the incipient to intermediate stages of weathering.
Figure 11: Normalised REE plots for the (a) upper and (b) lower flow of the Chhindwara profile and (c) the Bidar laterite. Note the change in scale between the two profiles. In each of the REE plots, the samples are normalised to the least-weathered (parent rock) samples (Table 4).

Fractionation of the lanthanides is evident in the changes in the slope and the abundance relative to 1 (parent rock) in the plots. Highlighted for emphasis are Ce and Eu.

Figure 12: Fractionation of Sm/Nd ratio as a function of the CIA during the incipient and intermediate stages of Deccan Traps basalt weathering.

Figure 13: The fractionation of Eu from the trivalent REE, expressed as the Eu/Eu* ratio, is anti-correlated with the (a) CIA and the (b) Rb/Sr ratio, indicating the loss of Eu is strongly linked to that of Sr, Ca, and Na (i.e. plagioclase dissolution) during basalt weathering. The Eu/Eu* ratio is calculated following normalisation to the parent rock values (ChQB12 for Chhindwara profile and BB1 for Bidar profile). A Eu/Eu* value of 1 (dashed line) represents no Eu anomaly between Sm and Gd relative to the parent rock. The alkali element enriched samples from the Chhindwara profile (very high Rb/Sr) and the Bidar samples are not shown on the Rb/Sr plot.
Laterite profiles with an allochthonous admixture
Lateritic weathering profiles

Laterite

Laterite profiles with an allochthonous admixture

lateritised ferricretes

Ferricrete profiles affected by in situ weathering
Ferricrete alteration profiles

Ferricrete

Autochthonous

100%

Allochthonous

100%
**Figure 3**

**a. BIDAR LATERITE PROFILE (BB)**

- **KEY CHARACTERISTICS**
  - Indurated laterite cap
  - Vermiform (tubular) laterite
  - Paleowatertable
  - Laterite comprising fretworks of Fe-rich segregations
  - Fe-rich segregations and agglomerations
  - Saprolite with increasing Fe-rich staining and segregations
  - Anorthite, Augite
  - Kaolinite, Goethite, Hematite
  - Pedogenetic clays

**b. CHHINDWARA PROFILE (ChQ)**

- **KEY CHARACTERISTICS**
  - Soil cover over upper ~20 cm
  - Samples 9a-9d through corestone centre to weathering rind exfoliates
  - Well-developed spheroidal weathering texture - corestone-rich horizon between ~1-3 m
  - Alteration increases downward as controlled by primary basalt structure (e.g., vesicularity, grain size, flow banding)
  - Chilled base of upper flow with laminated fabric
  - Undulose, brecciated flow top
  - Variably vesiculated (zeolite-filled amygdales) - Vesicle % increases toward flow top
  - Uniform weathering texture (no internal structure)
  - Plagioclase-phyric massive flow (unweathered basalt not exposed at base)
Figure 4

Upper flow (ChQB)

Lower flow (ChQA)

Figure 4a and Figure 4b show different aspects of the rock formations. Figure 4a highlights the upper flow (ChQB) section, while Figure 4b focuses on the lower flow (ChQA) section. The images provide detailed views of the geological structures and layers present in these areas.
Figure 5

**a. A–CN–K plot**

**b. SAF plot**

**LEGEND**
- Bidar profile
- Chhindwara lower flow (ChQA)
- Chhindwara upper flow (ChQB)
- Poladpur & Ambenali formations
- Sub-Recent-Tertiary basalt weathering profiles

**Chemical index of alteration (CIA)**

**Index of lateritisation (IOL)**

- Strongly lateritised
- Moderately lateritised
- Weakly lateritised
- Kaolonitised

**Al	extsubscript{2}O	extsubscript{3}, CaO*, Na	extsubscript{2}O, K	extsubscript{2}O, SiO	extsubscript{2}, Fe	extsubscript{2}O	extsubscript{3}**
Figure 6

Mafic index of alteration [MIA]

Chemical index of alteration

30 40 50 60 70 80 90 100

2+  3+

3.0

2.5

2.0

1.5

1.0

0.5

0.0

30 40 50 60 70 80 90 100

Chemical index of alteration
Figure 7

Mafic index of alteration (MIA)

1. $\text{MIA}^\text{R} = \frac{\text{Al}_2\text{O}_3}{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(\text{I}) + \text{MgO} + \text{CaO}^* + \text{Na}_2\text{O} + \text{K}_2\text{O}}$

2. $\text{MIA}^\text{O} = \frac{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(\text{II})}{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3(\text{II}) + \text{MgO} + \text{CaO}^* + \text{Na}_2\text{O} + \text{K}_2\text{O}}$

LEGEND
- Bidar profile
- Chhindwara lower flow (ChQA)
- Chhindwara upper flow (ChQB)
- Poladpur & Ambenali formations
- Sub-Recent-Tertiary mafic weathering profiles
- Predicted weathering trends
Figure 8

The figure illustrates the relationship between the chemical index of alteration (CIA) and the index of lateritisation (IOL) for different exposure times. The data points are color-coded to represent different stages of mineral weathering:

- **Primary Mineral Weathering and Early Weathering Products**: Represented by yellow squares.
- **Weathering Products Dominate**: Represented by red squares.
- **Most Resistant Minerals**: Represented by green squares.

The exposure times are indicated as:
- **20 x 10^3 years** < Exposure Time < **60 x 10^6 years**

The graph shows a trend where the CIA and IOL values change with exposure time, indicating the progression of weathering processes.
Figure 9

Weathering index

\( \tau_{Nb, j} \)
Figure 10

Chemical index of alteration

Element (j)
- Na
- Ca
- Mg
- Sr
- Ba
- Be
- Li
Figure 11

a. CHHINDWARA PROFILE (Upper flow - ChQB)

b. CHHINDWARA PROFILE (Lower flow - ChQA)

c. BIDAR PROFILE
Figure 12

Chemical index of alteration (CIA)

Sm/Nd
Figure 13
Table 1: Major\(^a\) (wt. %) and trace element (ppb) data for the Bidar (BB) laterite profile

<table>
<thead>
<tr>
<th>Depth (cm)</th>
<th>BB1</th>
<th>BB2</th>
<th>BB3</th>
<th>BB4</th>
<th>BB5</th>
<th>BB6</th>
<th>BB7</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO(_2)</td>
<td>48.9</td>
<td>50.06</td>
<td>38.59</td>
<td>38.78</td>
<td>30.61</td>
<td>6.12</td>
<td>36.68</td>
</tr>
<tr>
<td>TiO(_2)</td>
<td>2.16</td>
<td>2.29</td>
<td>5.11</td>
<td>4.78</td>
<td>5.76</td>
<td>1.4</td>
<td>2.44</td>
</tr>
<tr>
<td>Al(_2)O(_3)</td>
<td>13.72</td>
<td>14.15</td>
<td>31.54</td>
<td>31.95</td>
<td>25.83</td>
<td>6.97</td>
<td>31.3</td>
</tr>
<tr>
<td>Fe(_2)O(<em>3)(</em>{3(T)})</td>
<td>13.4</td>
<td>12.63</td>
<td>24.1</td>
<td>21.64</td>
<td>36.95</td>
<td>84.81</td>
<td>27.7</td>
</tr>
<tr>
<td>MgO</td>
<td>6.93</td>
<td>5.99</td>
<td>0.4</td>
<td>0.38</td>
<td>0.23</td>
<td>0.14</td>
<td>0.26</td>
</tr>
<tr>
<td>MnO</td>
<td>0.19</td>
<td>0.22</td>
<td>0.11</td>
<td>0.06</td>
<td>0.06</td>
<td>0.03</td>
<td>0.33</td>
</tr>
<tr>
<td>CaO</td>
<td>10.99</td>
<td>11.45</td>
<td>0.19</td>
<td>1.91</td>
<td>0.07</td>
<td>0</td>
<td>0.05</td>
</tr>
<tr>
<td>Na(_2)O</td>
<td>2.46</td>
<td>2.78</td>
<td>bdl</td>
<td>bdl</td>
<td>bdl</td>
<td>bdl</td>
<td>bdl</td>
</tr>
<tr>
<td>K(_2)O</td>
<td>0.16</td>
<td>0.25</td>
<td>0.02</td>
<td>0.01</td>
<td>0.02</td>
<td>0</td>
<td>0.03</td>
</tr>
<tr>
<td>P(_2)O(_5)</td>
<td>0.16</td>
<td>0.19</td>
<td>0.18</td>
<td>0.03</td>
<td>0.08</td>
<td>0.33</td>
<td>0.07</td>
</tr>
<tr>
<td>LOI</td>
<td>0.6</td>
<td>0.72</td>
<td>11.7</td>
<td>12.4</td>
<td>11.1</td>
<td>11.1</td>
<td>11.4</td>
</tr>
<tr>
<td>Total</td>
<td>99.07</td>
<td>100.01</td>
<td>100.24</td>
<td>99.54</td>
<td>99.61</td>
<td>99.8</td>
<td>98.86</td>
</tr>
</tbody>
</table>

CIA \(=\) 37 36 99 90 99 100 100
MIA\(_{\text{O}}\) \(=\) 35 36 97 91 99 99 98
IOL \(=\) 36 35 59 58 67 94 62

| Nb   | 9262 | 10202| 18943| 18600| 23480| 5646 | 14495|
| Li   | 4466 | 5159 | 7019 | 3736 | 4271 | 1202 | 13235|
| Rb   | 956  | 5678 | 1165 | 335  | 811  | 321  | 1524 |
| Cs   | 8    | 221  | 114  | 68   | 102  | 37   | 142  |
| Tl   | 3    | 24   | 8    | 16   | 29   | 7    | 272  |
| Be   | 606  | 632  | 2435 | 1289 | 1319 | 3814 | 859  |
| Sr   | 208492|227625|13746 |23686 |6022 |46072 |15494 |
| Ba   | 50844| 92866| 59309|10603 |16262|24813 |398559|
| La   | 8587 | 10314| 31316|4642  |9357 |53766 |26386 |
| Ce   | 21805| 24700| 25511|15509 |21649|119190|285003|
| Pr   | 3250 | 3729 | 16136|1724  |4044 |8776 |4007 |
| Nd   | 15508| 17765| 74513|7240  |17296|32632|11510 |
| Sm   | 4448 | 5035 | 28324|1717  |4166 |7276 |1901 |
| Eu   | 1592 | 1786 | 12010|490   |1134 |1832 |463  |
| Gd   | 5264 | 6039 | 62888|1657  |3688 |5340 |1444 |
| Tb   | 850  | 968  | 12625|288   |620  |859  |274  |
| Dy   | 5083 | 5758 | 91424|1791  |3642 |4878 |1516 |
| Ho   | 1028 | 1170 | 21863|365   |712  |926  |300  |
| Er   | 2688 | 3047 | 63014|1022  |1956 |2614 |848  |
| Tm   | 379  | 425  | 9016 |162   |310  |440  |137  |
| Yb   | 2290 | 2556 | 53585|1087  |2003 |3089 |933  |
| Lu   | 321  | 360  | 8079 |155   |280  |425  |131  |

\(^a\) Major element data has been recalculated without the LOI as reported in Borger and Widdowson (200
<table>
<thead>
<tr>
<th>Depth (cm)</th>
<th>Tb</th>
<th>Dy</th>
<th>Lu</th>
<th>Er</th>
<th>Gd</th>
<th>TbO₂</th>
<th>TiO₂</th>
<th>Mo</th>
<th>Nb</th>
<th>Nb₂O₃</th>
<th>NiO</th>
<th>Na₂O</th>
<th>MgO</th>
<th>CaO</th>
<th>P₂O₅</th>
<th>CO₂</th>
<th>SO₂</th>
<th>CH₄</th>
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<tbody>
<tr>
<td>2</td>
<td>7.36</td>
<td>7.45</td>
<td>7.42</td>
<td>7.43</td>
<td>7.41</td>
<td>7.42</td>
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<td>7.46</td>
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<td>7.49</td>
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<td>7.51</td>
<td>7.52</td>
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<td>7.46</td>
<td>7.43</td>
<td>7.44</td>
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<td>7.50</td>
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<td>7.59</td>
<td>7.60</td>
</tr>
</tbody>
</table>

Table 2: Major (wt. %) and trace element (ppb) data for the Chhindwara (ChQA) weathering profile

<table>
<thead>
<tr>
<th>Lower flow (ChQA)</th>
<th>Upp</th>
<th>ChQA1</th>
<th>ChQA2</th>
<th>ChQA3</th>
<th>ChQA4</th>
<th>ChQA5</th>
<th>ChQA6</th>
<th>ChQA7</th>
</tr>
</thead>
<tbody>
<tr>
<td>630</td>
<td>610</td>
<td>590</td>
<td>580</td>
<td>570</td>
<td>560</td>
<td>550</td>
<td>540</td>
<td>530</td>
</tr>
<tr>
<td>490</td>
<td>480</td>
<td>470</td>
<td>460</td>
<td>450</td>
<td>440</td>
<td>430</td>
<td>420</td>
<td>410</td>
</tr>
</tbody>
</table>

Table 2: Continued
Table 3: Mole conversions for weathering index calculations

moles Al₂O₃ = wt % Al₂O₃ ÷ 101.96
moles Fe₂O₃ = wt % *Fe₂O₃ ÷ 159.69
    moles FeO = wt % FeO ÷ 71.84
moles MgO = wt % MgO ÷ 40.30
moles CaO = wt % CaO ÷ 56.08
moles Na₂O = wt % Na₂O ÷ 61.98
moles K₂O = wt % K₂O ÷ 94.20
moles P₂O₅ = wt % P₂O₅ ÷ 141.94
moles CO₂ = wt % CO₂ ÷ 44.01

CaO* = moles CaO - moles CO₂ (calcite) - (0.5 x moles CO₂) (dolomite) - [(10/3) x moles P₂O₅] (apatite)

*wt % Fe₂O₃ = wt % FeO wt ÷ 0.8998

Click here to download Table: Table3_MoleConversions.xlsx
Table 4: Weathering index values for mafic standards and parent rock samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>CIA</th>
<th>MIA(_{(0)})</th>
<th>IOL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>USGS(^{a})</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BIR-1</td>
<td>36.3</td>
<td>30.5</td>
<td>35.8</td>
</tr>
<tr>
<td>BHVO-2</td>
<td>35.7</td>
<td>33.4</td>
<td>34.1</td>
</tr>
<tr>
<td>BCR-2</td>
<td>41.2</td>
<td>44.1</td>
<td>33.5</td>
</tr>
<tr>
<td>W-2</td>
<td>39.5</td>
<td>36.0</td>
<td>33.3</td>
</tr>
<tr>
<td>DNC-1</td>
<td>43.2</td>
<td>33.2</td>
<td>37.5</td>
</tr>
<tr>
<td><strong>Parent rocks</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BB-1</td>
<td>36.5</td>
<td>35.0</td>
<td>35.7</td>
</tr>
<tr>
<td>ChQB12</td>
<td>36.0</td>
<td>38.0</td>
<td>36.5</td>
</tr>
<tr>
<td>ChQB9d (corestone centre)</td>
<td>38.0</td>
<td>38.8</td>
<td>35.7</td>
</tr>
</tbody>
</table>

**Deccan Traps (Widdowson et al., 2000)**

|                        |      |               |      |
| average Poladpur formation | 37.8 | 37.1          | 35.9 |
| average Ambenali formation  | 37.2 | 37.4          | 37.2 |

\(^{a}\) Weathering index values were calculated using the accepted values for the USGS standards.
Click here to download Background dataset for online publication only: SupplementaryTable1.xlsx