The Loongana (CL) group of carbonaceous chondrites

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Abstract

A coordinated study of the petrology, mineral chemistry, and bulk chemical and isotopic composition of the five ungrouped carbonaceous chondrites Coolidge, Loongana 001, Los Vientos (LoV) 051, Northwest Africa (NWA) 033, and NWA 13400 reveals that these meteorites have a similar set of properties that distinguishes them from the other carbonaceous chondrite groups and allows definition of the new Loongana (CL) group of carbonaceous chondrites. The basic characteristics of the investigated samples are: (1) Lithophile element ratios (e.g., Al/Mg, Si/Mg) are within the typical range of other carbonaceous chondrite groups. (2) Fe-Ni metal abundances are considerably higher than for CV, but similar to CR chondrites. (3) Chondrule size-frequency distributions are similar to CV, but dissimilar to CR chondrites. (4) The mean CAI abundance is ~1.4 vol%, i.e., lower than in CV but much higher than in CR chondrites. (5) Very low amounts of matrix (17–21 vol%), the lowest among the main carbonaceous chondrite groups (CI, CM, CO, CV, CR, CK). (6) Olivine is nearly equilibrated, with mean fayalite (Fa) values between 12.5 mol% (Loongana 001) and 14.7 mol% (NWA 13400) as a metamorphic effect. (7) Lower Al2O3 and higher MgO and Cr2O3 concentrations in matrix, compared to matrix in CV, CK, and CR chondrites. (8) Volatile elements (Mn, Na, K, Rb, Cs, Zn, Se, Te, Pb, Tl) are considerably depleted compared to all other main carbonaceous chondrite groups, reflecting the low matrix abundance. (9) Bulk O isotope compositions plot along the CCAM line (\(\Delta^{17}\)O = –3.96 to –5.47‰), partly overlapping with the CV and CK chondrite fields but including samples that are more \(^{16}\)O-rich. (10) Unique positions of CL values in the \(^{54}\)Cr–\(^{50}\)Ti isotope plot, with \(^{54}\)Cr values similar to CV, CK, and CO, but \(^{50}\)Ti values similar to CR chondrites. All CL chondrites studied here are of petrologic type 3.9 to 4, indicating that they have been thermally metamorphosed on the parent body. The diagnostic features of CL chondrites detailed here provide a basis for identifying CL members of lower petrologic types. Such samples will be important for determining the pristine state of these meteorites and their components.

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1. INTRODUCTION

Chondrites are cosmic sediments, which consist of up to 92% chondrules (e.g. Weisberg et al., 2006; Metzler, 2012), i.e., once molten silicate spherules with sizes mostly between 0.1 and 1 millimeter. Further components are metal and sulfide grains, refractory objects like Ca-Al-rich inclusions and amoeboid olivine aggregates (CAIs, AOAs; e.g. Krot, 2019) and fine-grained matrix. These components occur in variable and often characteristic amounts in the distinct chondrite groups (e.g. Weisberg et al., 2006). Beside the Rumuruti (R) and Kakangari (K) groups, chondrites are subdivided into three classes, namely the ordinary, enstatite, and carbonaceous chondrites. The latter class consists of eight groups, namely the Ivuna-type (CI), Mighei-type (CM), Ormans-type (CO), Vigarano-type (CV), Karoonda-type (CK), Renazzo-type (CR), Bencubbin-type (CB), and CH (high iron) carbonaceous chondrites. Recently, King et al. (2019) defined the Yamato (CY) group of carbonaceous chondrites, which is not well established yet. Of these, the CI group is particularly distinctive, as members of this group do not contain any recognizable chondrules and because their bulk chemistry agrees closely with the heavy element component of the solar photosphere (e.g. Lodders, 2003; Lodders et al., 2009). Each chondrite group is thought to represent a single parent body (or several related parent bodies; e.g.,Gattacceca et al., 2020). As such, identification of new chondrite groups is important for understanding the diversity of planetesimals formed in the early solar system, and the processes that led to this diversity.

In addition to the established carbonaceous chondrite groups there are also ungrouped carbonaceous chondrites, which cannot be assigned to one of these groups. Examples are Tagish Lake, Tarda, Essebi and Bells (C2-ungrouped), as well as Ningqiang (C3-ungrouped). Based on similar properties some ungrouped chondrites can be combined into grouplets (consisting of less than 5 individual members), as in the case of Coolidge and Loongana 001 (Kalemeyn and Rubin, 1995). Coolidge is a well-studied meteorite which was originally classified as a CV4 chondrite (Van Schmus, 1969). However, McSween (1977a) concluded that it cannot simply be a metamorphosed sample of a reduced CV chondrite. By contrast, Kalemeyn and Wasson (1982) stated that Coolidge represents a CV chondrite, which “was probably subjected to open-system metamorphic reheating”, a conclusion also reached by Scott and Taylor (1985). However, Kalemeyn (1987) subsequently argued that the strong depletion of volatile elements more likely reflects nebular processes and that this meteorite, therefore, may not be a CV chondrite. Further, Noguchi (1994) found that, compared to the reduced CV chondrite Efremovka, chondrules and matrix in Coolidge are systematically depleted in Na, which also points to a nebular rather than a metamorphic process. Finally, based on petrographic observations and characteristic differences in the inventories of volatile elements, Kalemeyn and Rubin (1995) demonstrated that Coolidge clearly does not belong to the CV group, but together with the newly discovered meteorite Loongana 001, defines a new carbonaceous chondrite grouplet.

During comparative preexamination we found that the three ungrouped carbonaceous chondrites Los Vientos (LoV) 051, Northwest Africa (NWA) 033 and NWA 13400 have similar petrologic, mineralogical and microchemical properties as the Coolidge-Loongana 001 grouplet. The purpose of this study is to verify if these five samples are also similar regarding their bulk chemical and isotopic composition, allowing to group them together into a newly defined group of carbonaceous chondrites.

2. SAMPLES

Five ungrouped carbonaceous chondrite finds, namely Coolidge (USA, found 1937), Loongana 001 (Australia, found 1990), LoV 051 (Chile, found 2010), NWA 033 (Northwest Africa, found 1999), and NWA 13400 (Northwest Africa, found 2016) are investigated in this study. Polished thin sections (PTS), polished epoxy mounts and polished massive samples (slices) of these chondrites were used for petrographic and microchemical characterization. Subsample designations and masses of analyzed material are given in Table 1. To assess data quality and accuracy, representative bulk powder samples of the Allende CV3 chondrite were analyzed together with those of the above specimens. For comparison, we also determined the matrix fraction of Allende (CV3oA), Vigarano (CV3oA), and NWA 7020 (CR2) as well as the chondrule size-frequency distributions in Allende and NWA 7020.

In the following, the meteorite Loongana 001 is mostly termed Loongana for short and we collectively refer to the five investigated meteorites as “CL chondrites” and “CL samples”. The reasoning for naming the proposed new group “CL” is given in Section 5.3.1.

3. METHODS

All samples, including those that were subsequently powdered, were investigated and documented by optical and scanning electron microscopy. Samples of each meteorite were powdered in an agate mortar for obtaining the bulk chemical and isotopic compositions by various techniques. Aliquots for all laboratories, with the exception of LA-ICP-MS (Sc, Se, Te), were taken from the same sample powder (Table 1). All chemical data are given in wt.% or ppm (μg/g), respectively.

3.1. Polarizing microscopy

Polished thin sections (PTS) of all samples were used to determine shock stages and weathering grades by polarizing microscopy. Shock stages were determined according to the instructions given in Stöffler et al. (2018); weathering grades have been determined according to Wlotzka (1993). The latter scheme was initially developed to characterize ordinary chondrites, but we adopt it here for the use in cases of metal-rich carbonaceous chondrites.
3.2. Scanning electron microscopy (SEM)

The scanning electron microscope (SEM; Jeol JSM-6610LV) of the Interdisciplinary Centre for Electron Microscopy and Microanalysis (ICEM; University of Münster) was used to characterize the sample textures and to identify mineral phases. All subsamples, including slices for subsequent destructive chemical analysis, were documented by preparing photomosaics (backscatter electron images; BSE) of their polished surfaces. BSE images were also used to identify the various meteorite components (chondrules, CAIs, amoeboid olivine aggregates (AOAs), matrix) and to determine modal abundances and apparent chondrule sizes. For element mapping we used the SEM (Zeiss Sigma 300 VP) at the Institut für Geologie und Mineralogie, University of Cologne.

3.2.1. Energy-dispersive X-ray (EDX) analysis

Mineral analyses were obtained using an energy-dispersive X-ray (EDX) analyzing system (INCA, Oxford Instruments); all data were automatically normalized to 100 wt.%. Samples and appropriate mineral standards were measured at a working distance of 10 mm and an acceleration voltage of 20 KV. The beam current constancy was controlled by a Faraday cup. The standard ZAF correction procedures were applied (Bence and Albee, 1968). Minerals of known composition (olivine, low-Ca pyroxene and plagioclase from the L6 chondrite Leedey; Feldstein et al., 2001) were measured repeatedly as reference materials to ensure the reproducibility of the EDX system. All mineral analyses were checked for stoichiometry. Since none of the chosen analysis methods in this study was able to measure the bulk chondrite S content of the samples, the

Fig. 1. Overall texture and lithologic components of the Loongana 001 meteorite as an example for the proposed CL group of carbonaceous chondrites (a: back-scatter electron image; b: phase map including color coding). Beside the omnipresent metal-rich porphyritic Type-I chondrules (PO, POP) with irregular shapes (1,6), the following components are visible: (2) Barred olivine (BO) chondrule; (3) Al-rich chondrule; (4) Chondrule fragment; (5) Irregularly shaped chondrule adhering to a larger one; (6) Chondrule with thick pyroxene rim (7) and olivine-rich core; (8) Ca-Al-rich inclusion (CAI).
The following procedure was used to obtain a rough estimate of bulk S concentrations. By using the lowest possible magnification (30x) and a duration of 40 seconds each, EDX analyses of 2–3 different PTS regions of each CL sample and Allende were recorded and averaged. The mean S concentration for Allende, obtained by this method, is 1.67 wt. %, compared to the literature value of 2.10 wt. % (Jarosewich et al., 1987). The obtained values for S concentrations in the other samples were multiplied by the resulting correction factor of 1.26.

### 3.2.2. Element mapping (phase maps)

We performed element mapping in order to produce phase maps of entire chondrite sections. For this application, the electron beam scanned the stationary sample surface over small areas (~500 x 400 µm, pixel size ~4 µm) with a dwell time of 10 ms per pixel. The accelerating voltage was set to 20 kV and the aperture diameter to 60 µm. The sample was then moved to a new center and the next map was rastered. The phase maps (Fig. 1; Electronic Annex) were then produced with the PHAPS application.

### Table 1

<table>
<thead>
<tr>
<th>Sample allocation, weight, and designation of the investigated (sub)samples of CL chondrites.</th>
<th>Coolidge</th>
<th>Loongana 001</th>
<th>LoV 051</th>
<th>NWA 033</th>
<th>NWA 13400</th>
</tr>
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<tbody>
<tr>
<td>Powder sample, total (mg)</td>
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<td>565</td>
<td>513</td>
<td>598</td>
<td>610</td>
</tr>
<tr>
<td><strong>ICP-MS</strong>&lt;br&gt;University of Münster, Inst. f. Planetologie</td>
<td>128</td>
<td>137</td>
<td>81</td>
<td>163</td>
<td>178</td>
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<td>Sample powder (mg)</td>
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<td>86</td>
<td>87</td>
<td>84</td>
<td>86</td>
</tr>
<tr>
<td><strong>Laser fluorination</strong>&lt;br&gt;Open University, Milton Keynes</td>
<td>10</td>
<td>10</td>
<td>15</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Sample powder (mg)</td>
<td>131</td>
<td>130</td>
<td>130</td>
<td>134</td>
<td>135</td>
</tr>
<tr>
<td><strong>XRF</strong>&lt;br&gt;University of Cologne</td>
<td>-K-397-a-1 (52)</td>
<td>-F-a-1 (54)</td>
<td>-MS-a-1 (76)</td>
<td>PL-00027 (112)</td>
<td>-MS-a-1 (361)</td>
</tr>
<tr>
<td><strong>SEM-EDX, Polarizing microscopy</strong>&lt;br&gt;University of Münster, Inst. f. Planetologie</td>
<td>-K-397-a (52)</td>
<td>-F-a (44)</td>
<td>-MS-a (76)</td>
<td>-MS-a (126)</td>
<td>-MS-a (152)</td>
</tr>
<tr>
<td>Polished thin section numbers (size; mm²)</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>


### Fig. 2

(a) Frequency distributions of fayalite (Fa) values in olivine and low-Ca pyroxene, respectively, in CL chondrites. Coolidge, Loongana, and NWA 033 show narrow Fa distributions. The samples LoV 051 and NWA 13400 show broader distributions, indicating a less intensive metamorphic overprint. The numbers of analyzed olivine and pyroxene grains are listed in Tables 4 and 5, respectively. (b) Frequency distributions of ferrosilite (Fs) values in olivine and low-Ca pyroxene, respectively, in CL chondrites.
These display every mineral phase in false color and allow their visual identification. We further used the phase maps and the software "Measure" (Datinf GmbH, Tübingen, Germany) to determine the modal abundances of CAIs.

3.2.3. Determination of apparent (2D) chondrule sizes

The distributions of apparent (2D) chondrule sizes in the five CL chondrites have been measured, plus those in Allende (CV3$_{oxA}$) and NWA 7020 (CR2) for comparison. For this purpose the software "Measure" (Datinf GmbH, Tübingen, Germany) was used, stringently following a raster on the BSE photomosaics to not overlook any chondrule cut face. Apparent chondrule sizes were obtained by measuring the diameters of chondrule cut faces at an angle of 45° from the line of the maximum apparent diameter, and thereby a reasonable average between minimum and maximum is obtained. It was calculated by Eisenhour (1996) that this method provides a good approximation (within 2%) of the diameter of a circle equal in area to that of the measured ellipse. Except for metal grains, sulfide grains, and CAIs, every subrounded, more or less isometric (aspect ratio <~2) object was considered as a chondrule. Chondrules consisting of welded smaller chondrules were measured as one single chondrule. The same holds for chondrules with adhering layers of smaller chondrules and for the rare chondrules with igneous rims. Chondrule fragments with angular fracture surfaces were only included when their original outline could be unequivocally reconstructed (~half chondrules). Other chondrule fragments and angular monomineralic grains were omitted. It cannot be ruled out that the data sets contain small fractions of unrecognized roundish chondrule fragments, especially in the smallest size bins (<300 μm). Distributions of apparent chondrule sizes (2D) were already measured by this method in several ordinary chondrites (Metzler, 2018; Metzler et al., 2019).

3.3. Electron microprobe (EMPA)

The electron microprobe (JEOL 8900RL) at the Institut für Geologie und Mineralogie (University of Cologne) was used for chemical analysis. Mineral compositions (olivine, low-Ca pyroxene, Ca-rich pyroxene) were determined by spot analyses using a focused, stationary beam of 1 nA. For the analyses of matrix, the beam was defocused along linear tracks (50 mm each) across the polished sample surfaces (thin and thick sections) was done using 193 nm ArF excimer laser (Analyte G2, Photon Machines). A repetition rate of 10 Hz and an energy of ~3–4 J/cm2 were used throughout the entire session. The beam spot diameter varied between 80 and 100 μm. Elemental analysis has been carried out with an Element 2 mass spectrometer (Thermo-Fisher). Forward power was 1300 W and reflected power <2 W. Gas flow rates were 1.2 l/min for He (carrier gas of ablated material), 0.8 l/min and 1.1 l/min for the Ar-auxiliary and sample gas, respectively. Cooling gas flow rate was set to 16 l/min. Before starting analysis, the system has been tuned (torch position, lenses, gas flows) on a NIST 612 glass, measuring $^{139}$La, $^{232}$Th and $^{232}$Th$^{16}$O to get stable signals and high sensitivity, as well as low oxide rates ($^{232}$Th$^{16}$O/$^{232}$Th < 0.1%) during ablation. The above elements were as described in Stracke et al. (2012) has been used, with a few additional, minor modifications. A sample to standard ratio of 1:30 was used, with typically 120 mg sample material mixed with 3.6 g lithium tetaborate (Li$_2$B$_4$O$_7$) flux agent. Before mixing, the sample was pre-treated with aqua regia for a few hours at 120 °C to oxidize the metals. A Claisse LeNeo fluxer was used to automatically produce 27 mm diameter glass disks. The Panalytical Zetium sequential wavelength dispersive X-ray spectrometer at the University of Cologne was used for sample measurements. Standard calibration lines were calculated from 7 identically prepared standard rock samples. The precision for all elements analyzed is below 1% (Wolf and Palme, 2001). An Allende glass disk was produced and measured in the same way as the samples. The obtained data for Allende are in excellent agreement with literature data. Some CL samples showed low totals which most likely resulted from loss of sample material (~2 mg, estimated) during sample preparation as well as loss of material during sample fluxing. The latter was possibly caused by the decomposition of weathering products (hydroxides, calcite) and loss of volatile components (H$_2$O, CO$_2$). There was only enough powder to determine the loss on ignition (LOI) of NWA 13400 (2.76%, when 120 mg are heated at 1000 °C for 2 h). This LOI equals ~3 mg loss of sample material. We recalculated samples with low totals considering an estimated total loss of material of 5 mg. However, some totals are still low, possibly indicating an underestimation of material loss.

3.5. Trace element compositions

Trace element concentrations were analyzed on aliquots of the initial representative meteorite powders and an Allende Standard powder (Allende MS-A). The elements Li, V, Co, Rh, Sr, Y, Zr, Nb, Mo, Cs, Ba, the REE, Hf, Ta, Ti, Pb, Th, and U were analyzed by isotope dilution sector field inductively coupled plasma mass spectrometry (ID-SF-ICP-MS), as described in detail in Stracke et al. (2012).

Concentrations of Sc, Se, and Te in the samples and Allende were analysed by LA-ICP-MS at the University of Münster (Institut für Mineralogie). Sample ablation along linear tracks (50 mm each) across the polished sample surfaces (thin and thick sections) was done using 193 nm ArF excimer laser (Analyte G2, Photon Machines). A repetition rate of 10 Hz and an energy of ~3–4 J/cm2 were used throughout the entire session. The beam spot diameter varied between 80 and 100 μm. Elemental analysis has been carried out with an Element 2 mass spectrometer (Thermo-Fisher). Forward power was 1300 W and reflected power <2 W. Gas flow rates were 1.2 l/min for He (carrier gas of ablated material), 0.8 l/min and 1.1 l/min for the Ar-auxiliary and sample gas, respectively. Cooling gas flow rate was set to 16 l/min. Before starting analysis, the system has been tuned (torch position, lenses, gas flows) on a NIST 612 glass, measuring $^{139}$La, $^{232}$Th and $^{232}$Th$^{16}$O to get stable signals and high sensitivity, as well as low oxide rates ($^{232}$Th$^{16}$O/$^{232}$Th < 0.1%) during ablation. The above elements were
quantitatively analysed using the NIST 612 glass as an external standard and 44Ca as internal standard; Ca concentrations had been previously determined by XRF. Concentrations of measured elements were calculated using the Glitter software (Griffin et al., 2008). Standard reference glasses GSD1-G and GSE1-G were analysed as monitor for precision and accuracy. Obtained results match the published range of concentrations given in the GeoReM database (version 24; Jochum et al., 2005).

### 3.6. Titanium and chromium isotopes

Aliquots (~35 mg each) of the initial representative meteorite powders (equivalent to ~20–30 μg Ti) were digested in HF-HNO3-HClO4 (2:1:0.05) on a hotplate at 180–200 °C for five days and in aqua regia (HCl-HNO3) at 130–150 °C for another two days. Thereafter, the samples were dissolved in 12 M HNO3, and ~65 mg H3BO3 was added. Ti was separated from the sample matrix via a two-stage anion exchange chromatography adapted from the previously established procedure from Zhang et al. (2011). In a first step, the sample solutions were loaded onto columns prepacked with 2 ml TOGDA® anion exchange resin, where Ti was eluted in 20 ml 12 M HNO3–1 wt.% H2O2. Afterwards, the Ti cuts were dissolved in 2.5 ml 4 M HF and loaded onto clean-up columns filled with 0.8 ml Bio-Rad® AG1-X8 anion exchange resin, where Ti was eluted in 6 ml 9 M HCl–0.01 M HF.

Titanium isotope measurements were performed in two lines using a ThermoScientific Triton Plus Thermal Ionization Mass Spectrometer (Institut für Planetologie, University of Münster) in static mode. The sample solutions (containing ~500 ppm Cr in 6 M HCl) were loaded on 4–6 filaments and each filament was measured multiple times, with total ion beam intensities of ~1.4 × 10^−10 A on 52Cr. Instrumental mass fractionation was corrected assuming a constant 50Cr/52Cr = 0.051859 and using the exponential law. The data are reported in ε54Cr values as the parts per ten thousand deviation from the terrestrial NIST SRM3112a Cr standard. The sample uncertainty is reported as the Student-t 95% confidence interval based on repeated analyses (N = 12) of the sample solution.

Chromium was collected during the first step of the two-stage anion exchange chromatography used for the separation of Ti, where Cr is eluted in 25 ml 12 M HNO3 (+trace H3BO3) together with most other matrix elements. Aliquots (equivalent to ~30 μg Cr) were taken from this solution, dried down, and redissolved in 1 ml 6 M HCl. Afterwards, Cr was separated from the sample matrix using a single-stage anion exchange chemistry for the removal of Fe (Bio-Rad® AG1-X8 anion exchange resin), followed by a two-stage cation exchange chromatography (Bio-Rad® AG50W-X8 cation exchange resin) as described by Yamakawa et al. (2009), including a four-day conversion of Cr[III]Cl3/Cr[II]Cl2 to Cr³⁺.

Chromium isotope measurements were performed using a ThermoScientific Triton Plus Thermal Ionization Mass Spectrometer (Institut für Planetologie, University of Münster) in static mode. The sample solutions (containing ~500 ppm Cr in 6 M HCl) were loaded on 4–6 filaments and each filament was measured multiple times, with total ion beam intensities of ~1.4 × 10^−10 A on 52Cr. Instrumental mass fractionation was corrected assuming a constant 50Cr/52Cr = 0.051859 and using the exponential law. The data are reported in ε54Cr values as the parts per ten thousand deviation from the terrestrial NIST SRM3112a Cr standard. The sample uncertainty is reported as the Student-t 95% confidence interval based on repeated analyses (N = 8–12).
Fig. 3. Chondrule textures in CL chondrites. (a) Typical porphyritic olivine-pyroxene (POP) chondrule with irregular and lobate outline (Coolidge). (b) Chondrule with several protuberances in NWA 13400. (c) Assortment of differently sized and irregularly shaped chondrules or melt shreds (NWA 13400). (d) Complex chondrule (white outline), consisting of a porphyritic olivine (PO) chondrule core with irregular layers of adhering smaller chondrules or melt shreds, stacked on top of each other. The central chondrule is separated from these layers by an Fe-Ni layer (white). (e) Layered olivine chondrule with concentric inner Fe-Ni layer and thin pyroxene rim (LoV 051). (f) Multi-layered olivine chondrule with two concentric inner Fe-Ni layers and an outer Fe-Ni layer (NWA 13400). (g) Al-rich chondrule with fine-grained quench texture and large Al-rich enstatite crystals (dark gray; Loongana). (h) Barred olivine (BO) chondrule of Type-II with igneous rim in NWA 13400. SEM-BSE images; dark gray: low-Ca pyroxene; medium gray: olivine; white to light gray: Fe-Ni metal, sulfides. White to light gray linear features are terrestrial oxidation products along cracks (SEM-BSE images).
3.7. Oxygen isotopes

Oxygen isotope analyses were performed at the Open University using an infrared laser-assisted fluorination system (Miller et al., 1999; Greenwood et al., 2017). Samples and standards were weighed out into a Ni sample block with each aliquot having a mass of approximately 2 mg. The block was loaded into a vacuum-tight chamber and heated to 70 °C under vacuum overnight to remove surface-bound moisture. The system was then flushed with BrF3 to reduce the final blank level to < 60 nmoles O2.

Surface-bound moisture. The system was then flushed with field of 1 T and a sensitivity of ing Sample Magnetometer (VSM) with a maximum applied component magnetization (MRS) to saturation magnetization susceptibility of hysteresis loops provided the ratio of saturation remanent coercive force (Bc). High field susceptibility (χHF) was determined by a linear fit for applied fields >0.9 T of the hysteresis loops. Remanent coercive force (Bc, remanent) was determined by back field experiments performed with the VSM. The low field specific susceptibility (χ in m3/kg) was measured using Agico MFK1 apparatus with sensitivity of 5 × 10⁻¹⁰ Am². The analysis of hysteresis loops provides the ratio of saturation remanent magnetization (MRS) to saturation magnetization (Ms) and the coercive force (Bc).

The samples NWA 033 and LoV 051 are listed as C3-ungrouped in the MBDB, without allocation to a certain petrologic subtype. NWA 13400 is listed as C3-ungrouped, with a proposed petrologic subtype of 3.9. One of the standard methods to identify a chondrite’s subtype is determining the frequency distribution of fayalite (Fa) values of randomly chosen olivine grains. The percentage mean deviation (PMD) of these values is used as a measure for the equilibration state of a given meteorite (e.g. Hutchison, 2004; his Table 2.6). We determined these distributions in all samples, together with the frequency distributions of ferrosilite (Fs) values of their low-Ca pyroxene grains; the results are shown in Fig. 2. Occasionally, chondrules and chondrule fragments with relict Mg-rich core compositions of olivine can be found in all samples. The

3.8. Magnetic measurements

All magnetic measurements were performed at CEREGE (Aix-en-Provence, France). Hysteresis measurements were performed with a Princeton Micromag Vibrating Sample Magnetometer (VSM) with a maximum applied field of 1 T and a sensitivity of ~5 × 10⁻⁹ Am². The analysis of hysteresis loops provided the ratio of remanent magnetization (MRS) to saturation magnetization (Ms) and the coercive force (Bc, remanent). High field susceptibility (χHF) was determined by a linear fit for applied fields >0.9 T of the hysteresis loops. Remanent coercive force (Bc, remanent) was determined by back field experiments performed with the VSM. The low field specific susceptibility (χ in m3/kg) was measured using Agico MFK1 apparatus with sensitivity of 5 × 10⁻¹⁰ Am², operating at 200 A/m and a frequency of 976 Hz. We determined the S-300 ratio, defined as the isothermal remanent magnetization (IRM) obtained after applying a 3 T field and then a back field of 0.3 T normalized to the IRM acquired in 3 T. For this experiment, IRM were imparted using a pulse magnetizer from Magnetic Measurement.

4. RESULTS

This study will show that the three ungrouped carbonaceous chondrites Los Vientos (LoV) 051, Northwest Africa (NWA) 033 and NWA 13400 have similar properties as the Coolidge-Loongana 001 grouplet. This makes it possible to combine these five samples into a new carbonaceous chondrite group, which we term the Loongana (CL) group (see Section 5.3.1). In the following we define the basic criteria for identifying further members of this group. Since CL chondrites have some similarities to CVred chondrites (e.g., McSween, 1977a; Gattacceca et al., 2020) and CR chondrites (e.g., Kallemeyn et al., 1994), our discussion concentrates on these groups when comparing petrographic, chemical and isotopic data of CL chondrites.

4.1. Meteorite textures

The investigated samples are mainly composed of the typical carbonaceous chondrite components, namely Type-I chondrules (FeO-poor, metal rich), refractory inclusions (CAIs, AOAs), inter-chondrule sulfide and metal grains and a fine-grained matrix. Some chondrules of the chemical Type-II occur, as well. Despite the thermal overprint of all investigated CL samples, their Type-I and Type-II chondrules are still discernable by texture and mineral composition (see Section 4.7.1.1). Cut faces of the least weathered sample (LoV 051) show well-preserved tiny metal beads within and at the periphery of chondrules. Occasionally, large fragments of chondrules and refractory inclusions occur, but all samples are macroscopically unbrecciated. Their overall textures show some resemblance to the reduced CV chondrites (CVred) and CR chondrites, but differ in having a very low amount of matrix (see Section 4.7.3) and consequently a relatively high chondrule abundance. The compositions and textures of Coolidge and Loongana have been described by Noguchi (1994) and Kalllemeyn and Rubin (1995). The overall texture of Loongana is shown in Fig. 1 both as a photomosaic of SEM-BSE images (a) and a phase map (b). The following components can be distinguished in Fig. 1: abundant porphyritic (1,6) and rare non-porphyritic chondrules (2), rare AI-rich chondrules (3), chondrule fragments (4), adhering chondrules (5), olivine chondrules (6) with thick pyroxene rims (7), and CAIs (8). These components are described in detail below (Section 4.7).

4.2. Petrologic types

Coolidge and Loongana are listed as being C4-ungrouped in the Meteoritical Bulletin Database (MBDB; www.lpi.usra.edu/meteor/). However, they have also been classified as type 3.8 to 4 and type 3.8 by Kallemeyn and Rubin (1995) and Guimon et al. (1995). The samples NWA 033 and LoV 051 are listed as C3-ungrouped in the MBDB, without allocation to a certain petrologic subtype. NWA 13400 is listed as C3-ungrouped, with a proposed petrologic subtype of 3.9. One of the standard methods to identify a chondrite’s subtype is determining the frequency distribution of fayalite (Fa) values of randomly chosen olivine grains. The percentage mean deviation (PMD) of these values is used as a measure for the equilibration state of a given meteorite (e.g. Hutchison, 2004; his Table 2.6). We determined these distributions in all samples, together with the frequency distributions of ferrosilite (Fs) values of their low-Ca pyroxene grains; the results are shown in Fig. 2. Occasionally, chondrules and chondrule fragments with relict Mg-rich core compositions of olivine can be found in all samples. The
PMD values for Coolidge, Loongana, and NWA 033 are below 5% (Table 2), indicating that these samples belong to petrologic type 4, whereas those of LoV 051 and NWA 13400 are between 5% and 10%, indicating a 3.9 subtype.

4.3. Shock stages

All olivine grains in the investigated CL chondrites show undulatory extinction, but grains with planar fractures are rare or absent. Thus, all these samples are weakly shocked (Table 2) and belong to shock stage C-S2 (Stöffler et al., 2018), corresponding to the previously defined shock stage S2 (Stöffler et al., 1991).

4.4. Weathering grades

Most samples are moderately to heavily weathered (W2-W3/4) and crosscut by veins composed of the secondary alteration products of metals and sulfides. Nevertheless, variable amounts of Fe-Ni metal are still visible on cut faces of all samples. The estimated degrees of weathering for the CL samples, which were determined using the scheme of Wlotzka (1993), are given in Table 2. The least weathered CL chondrite of this study is LoV 051 (W1/2), which still contains considerable amounts of pristine metal.

4.5. Magnetic properties

The magnetic properties of all CL chondrites from this study are given in Table 3, together with data for the various CV subgroups (CV red, CV oxA, CV oxB) and CR chondrites for comparison. The magnetic properties of all samples, with the notable exception of LoV 051, have been considerably modified by terrestrial weathering. Thus, the data for LoV 051 are the most relevant for defining the primary characteristics of the proposed CL group. Its magnetic susceptibility (Table 3) is \( \log \chi = 5.24 \) (where \( \chi \) is in \( 10^{-7} \text{ m}^3 \text{ kg}^{-1} \)), higher than the corresponding values for the other samples. This difference can be accounted for by terrestrial weathering, which lowers the susceptibility of metal-bearing meteorites (Rochette et al., 2008). The susceptibility of LoV 051 (\( \log \chi = 5.24 \)) is slightly higher compared to CR chondrites, which have an average \( \log \chi = 5.04 \pm 0.12 \) as measured on 14 CR meteorites (Rochette et al., 2008), and almost identical to the value for the only CR chondrite fall Renazzo (\( \log \chi = 5.17 \)). The observed hysteresis loop for LoV 051 is typical of kamacite-bearing rocks (Gattacceca et al., 2014), including a curvature up to above 700 mT, very low coercivity (\( B_C = 3.08 \text{ mT} \)), very low \( M_{RMS}/M_S \) ratio (0.0155), and high \( B_{CR}/B_C \) ratio (6.10). Together with the \( S_{300} \) value of 1.00, there is strong evidence that the magnetic mineralogy is dominated by kamacite.

4.6. Modal compositions

The modal compositions of all CL chondrites from this study are summarized in Table 3.
2. The modal abundances (area% = vol%) of chondrules and chondrule fragments in our samples vary between 67 vol% (Loongana) and 79 vol% (Coolidge). The inter-chondrule metal and sulfide concentration, modified by terrestrial weathering, varies between 3 vol% (Coolidge) and 6 vol% (NWA 033, NWA 13400; Table 2). The concentration of CAIs, measured on phase maps (e.g., Fig. 1), appears highly variable (from 0.1 vol% for Coolidge to 8 vol% for Loongana). For the modal abundance of matrix we obtained values between 17 vol% (LoV 051, NWA 13400) and 21 vol% (NWA 033; Table 2).

To verify our method, we also determined the abundance of matrix in Allende (CV oxA) and Vigarano (CV red) and found values of 46 vol% and 35 vol%, respectively. The value for Allende is in the range of values obtained by McSween (1977a; 38.4 vol%) and Gattacceca et al. (2020; 50 vol%). Our Vigarano value is nearly identical to that of McSween (1977a; 34.5 vol%) and identical to that given in Gattacceca et al. (2020). Thus, our method reproduces literature values well, demonstrating that CL chondrites contain significantly less matrix than CV chondrites and most other carbonaceous chondrites.

4.7. Lithologic components

4.7.1. Chondrules

The investigated CL chondrites consist mainly of chondrules (67–79 vol%; Table 2), including minor amounts of chondrule fragments. In the following the term “chondrules” also includes chondrule fragments, if not stated otherwise.

4.7.1.1. Chondrule textures, types, and mineralogical compositions. Chondrules in the investigated meteorites are mostly porphyritic olivine (PO) and porphyritic olivine-pyroxene (POP) chondrules of Type-I. A high percentage of these show pyroxene rims (see Section 4.7.1.3). Many of these have non-circular, irregular boundaries, which has already been described for Coolidge and Loongana by Kalleney and Rubin (1995). Many chondrules in our samples, especially the smaller ones, give the impression of rapidly crystallized droplets from fragmented or shredded melts, which partly accumulated in a viscous state. They show remarkably irregular shapes with lobate outlines and protuberances (Fig. 3a) and sometimes display an amoeboidal shape (Fig. 3b). In some cases it is difficult to define the outlines and boundaries between adjacent objects (Fig. 3c). Many chondrules are the result of adhesive growth from smaller chondrules, i.e., they are part of welded chondrule agglomerates or adhere to the surface of larger chondrules. They form irregular layers of smaller roundish chondrules stacked on top of each other. A striking example is shown in Fig. 3d (white outline), which appears to be an agglomeration of small chondrules and lobate melt droplets around a central chondrule, separated by an Fe-Ni layer (white).

Fig. 4. Opaque phases in CL chondrites. (a) Polycrystalline kamacite grain in a Coolidge chondrule, showing taenite exsolutions (light gray) and exsolutions of chromite and merrillite (black dots) along kamacite grain boundaries; (b) Rare side-by-side occurrence of isolated kamacite (right center) and taenite grains (left center) in a chondrule in NWA 033; other grains are FeS; (c) Intergrowth texture of sulfide (FeS; gray) and kamacite (white) in a LoV chondrule; (d) Very large polycrystalline FeS grain in a Coolidge chondrule; white: kamacite. (SEM-BSE images).
Chondrule olivine is nearly equilibrated in most samples (Table 4; Fig. 2a). The lowest Fa values for olivine grains found in this study (beside the rare relict Mg-rich olivine cores) are around 10 mol%. This value is also the arbitrarily defined boundary between the chondrule Type-I (<10 mol% Fa) and Type-II (McSween, 1977b; Jones et al., 2005; Lauretta et al., 2006). However, the presence of Fe-Ni metal and low-Ca pyroxene with Fs values considerably below 10 mol% in most chondrules in our samples (Fig. 3a-f) relates them to Type-I. In all samples Mg-rich relict olivine cores are found, a further proof that Type-I chondrules were present prior to thermal metamorphism. Occasionally, chondrules of chemical Type-II can be observed in the samples (e.g. BO chondrule in Fig. 3h), identifiable by, e.g., their metal-free composition and the absence of pyroxene rims.

In case of larger chondrules there is a tendency that metal occurs as outer shells (e.g., Fig. 3g). In several cases, metal forms one or two concentric layers within the chondrules (Fig. 3e and f), which may indicate that these structures formed by agglomeratic growth. The largest Fe-Ni metal grains in chondrules were found in LoV 051 (1.1 mm) and Coolidge (1.6 mm). The metal mainly consists of polycrystalline kamacite with small amounts of taenite exsolutions (Fig. 4a). The Ni concentration in kamacite is fairly constant in all samples, with mean concentrations ranging between 7.1 wt.% (NWA 13400) and 8.0 wt.% (NWA 033). Measured Ni concentrations in taenite are variable, with values between 13.5 wt.% and 41.0 wt.%. This is consistent with the results of Scott and Taylor (1985) and Noguchi (1994) who found mean Ni concentrations of 6.4–7.1 wt.% and 19.2–20.9 wt.% for kamacite and taenite, respectively, in Coolidge. In many kamacite grains, tiny subrounded grains of chromite, merrillite, and SiO₂ occur as exsolutions along internal metal grain boundaries (Fig. 4a). A side-by-side occurrence of isolated kamacite and taenite grains in chondrules was only rarely observed (Fig. 4b). Polycrystalline, primary sulfide grains (troilite, FeS) up to 500 μm large and frequently intergrown with kamacite, can also be found in these chondrules (Fig. 4c).

Table 4
Mean chemical composition of olivine in the investigated CL chondrites (EMPA; wt.%).

<table>
<thead>
<tr>
<th></th>
<th>Coolidge</th>
<th>Loongana 001</th>
<th>LoV 051</th>
<th>NWA 033</th>
<th>NWA 13400</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>40.9 ± 0.2</td>
<td>41.1 ± 0.3</td>
<td>40.5 ± 0.3</td>
<td>40.7 ± 0.2</td>
<td>40.2 ± 0.4</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.07 ± 0.05</td>
<td>0.06 ± 0.04</td>
<td>0.04 ± 0.04</td>
<td>0.05 ± 0.03</td>
<td>0.03 ± 0.03</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>0.05 ± 0.08</td>
<td>0.02 ± 0.04</td>
<td>0.02 ± 0.02</td>
<td>0.03 ± 0.04</td>
<td>0.03 ± 0.05</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.09 ± 0.18</td>
<td>0.04 ± 0.03</td>
<td>0.05 ± 0.04</td>
<td>0.04 ± 0.03</td>
<td>0.05 ± 0.05</td>
</tr>
<tr>
<td>FeO</td>
<td>13.8 ± 0.4</td>
<td>11.9 ± 0.4</td>
<td>12.5 ± 1.1</td>
<td>12.8 ± 0.6</td>
<td>14.1 ± 1.2</td>
</tr>
<tr>
<td>MgO</td>
<td>45.8 ± 0.3</td>
<td>47.0 ± 0.4</td>
<td>47.5 ± 0.8</td>
<td>46.2 ± 0.5</td>
<td>45.8 ± 1.1</td>
</tr>
<tr>
<td>CaO</td>
<td>0.03 ± 0.04</td>
<td>0.02 ± 0.02</td>
<td>0.01 ± 0.01</td>
<td>0.02 ± 0.02</td>
<td>0.04 ± 0.04</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.01 ± 0.01</td>
<td>0.01 ± 0.02</td>
<td>0.01 ± 0.02</td>
<td>0.01 ± 0.02</td>
<td>0.01 ± 0.02</td>
</tr>
<tr>
<td>NiO</td>
<td>0.02 ± 0.02</td>
<td>0.02 ± 0.02</td>
<td>0.02 ± 0.02</td>
<td>0.02 ± 0.02</td>
<td>0.03 ± 0.03</td>
</tr>
<tr>
<td>Total</td>
<td>101.00</td>
<td>100.40</td>
<td>100.89</td>
<td>100.11</td>
<td>100.52</td>
</tr>
<tr>
<td>Fa</td>
<td>14.5 ± 0.4 (12.8–15.1)</td>
<td>12.5 ± 0.4 (11.0–13.2)</td>
<td>12.8 ± 1.2 (10.5–15.0)</td>
<td>13.5 ± 0.6 (10.1–14.5)</td>
<td>14.7 ± 1.3 (10.2–16.0)</td>
</tr>
<tr>
<td>FeO/MnO</td>
<td>60</td>
<td>52</td>
<td>52</td>
<td>53</td>
<td>61</td>
</tr>
<tr>
<td>n</td>
<td>44</td>
<td>49</td>
<td>47</td>
<td>46</td>
<td>30</td>
</tr>
</tbody>
</table>

Fa: Fayalite content in olivine (mol%); ±: ±1 standard deviation; Numbers in parenthesis: range of measured values.

Fig. 5. Frequency distributions of apparent chondrule sizes (100 μm binning), measured on BSE photomosaics. (a) Frequency distributions in the investigated CL chondrites; (b) Average distribution in CL chondrites (blue solid line, calculated from data in (a). Frequency distributions for the CV chondrites Allende (CV₉₋₁₀; red dashed line) and the CR chondrite NWA 7020 (black dotted line), obtained in this study, are shown for comparison. Please note the different scaling of the size axes in (a) and (b). n: number of measured chondrules.
and d). In some chondrule cut faces the sulfide + metal portion exceeds 50 area%. Some metal grains up to 1 mm in size occur outside of chondrules. Chondrule plagioclase compositions are variable (An72-An99) with negligible K concentrations. Noguchi (1994) found values of An 27-An95 in Coolidge. Plagioclase is concentrated in chondrule mesostases, frequently in the shape of lath-like intergrowths with Ca-rich pyroxene.

Some Al-rich chondrules have been observed in Loongana (Fig. 3g) and LoV 051, and non-porphyritic chondrules (barred olivine; BO) were found in Coolidge and NWA 13400 (Fig. 3h).

4.7.1.2. Chemistry of main minerals. The chemical compositions of the main mineral phases found in chondrules (olivine, low-Ca pyroxene, and Ca-rich pyroxene) were measured by EMPA. Relict Mg-rich olivine cores are occasionally found in all samples, but were not measured during the random analyses. A number of 30 to 49 olivine grains were measured in each sample and the variation of Fa values is shown in Fig. 2a. The broad distributions of LoV 051 and NWA 13400 in this figure mirror the lower grade of metamorphism of these meteorites (petrologic type 3.9; Table 2), compared to the other samples (petrologic type 4; Table 2). The distribution widths for LoV 051 and

---

Fig. 6. Textures and components of refractory inclusions in CL chondrites. (a) Complex CAI in Loongana, consisting of fine-grained subunits (anorthite, spinel) which are rimmed by Ca-rich pyroxene (medium gray), and some FeS (white); (b) CAI in NWA 13400, mainly consisting of subhedral-euhedral spinel (medium-dark gray), FeS (white), partly surrounded by a spinel rim; (c) CAI in Coolidge, rimmed by a spinel and diopside layer, probably a former Wark-Lovering rim. This inclusion contains a mineral assemblage consisting of FeS, TiO2, and spinel, embedded in extremely fine-grained groundmass; (d) Cl-apatite in a CAI in NWA 13400 (left center, light gray); (e) Refractory metal nugget (light gray to white; rich in Os, Ir, Pt, Ru) in a CAI in Loongana; (f) Amoeboid olivine aggregate (AOA) in LoV, intermingled with CAI material (right part of the inclusion). White to light-gray linear features in the images are terrestrial oxidation products along cracks (SEM-BSE images).
NWA 13400 are similar, but the distribution for the latter appears more asymmetric. This is probably a statistics problem, since 30 olivine grains were analyzed in NWA 13400, compared to 47 grains in LoV 051. The mean Fa values (mol%; Table 4) vary between 12.5 (Loongana) and 14.7 (NWA 13400). The Fa values for Coolidge and Loongana (14.5 and 12.5) are similar to those presented in earlier studies. Scott and Taylor (1985) and Noguchi (1994) found Fa values of 14 in Coolidge; Kallemeyn and Rubin (1995) give a value of 13.6. The mean Fa value found by the latter is 11.8. The mean FeO/MnO ratios (wt.%/wt.%) in olivine of our samples vary between 52 (Loongana) and 61 (NWA 13400). Noguchi (1994) found a mean value of 64 in Coolidge, compared to our value of 60.

Between 20 and 49 low-Ca pyroxene grains were measured in each sample and the Fs variation is presented in Fig. 2b. The mean Fs values (mol%; Table 5) vary between 6.6 (NWA 13400) and 10.4 (Loongana). Noguchi (1994) and Kallemeyn and Rubin (1995) found mean Fs values of 10.1 and 9.3, respectively, in Coolidge, compared to our value of 9.4. The mean Fs value of 6.9 for Loongana given in Kallemeyn and Rubin (1995) is somewhat lower than our value (10.4). The mean wollastonite (Wo) values in all samples vary between 0.9 and 1.1 mol% (Table 5). The mean Fs value of 6.9 for Loongana given in Kallemeyn and Rubin (1995) is somewhat lower than our value (10.4). The mean wollastonite (Wo) values in all samples vary between 0.9 and 1.1 mol% (Table 5).

### Table 5
Mean chemical composition of low-Ca pyroxene in the investigated CL chondrites (EMPA; wt.%).

<table>
<thead>
<tr>
<th></th>
<th>Coolidge</th>
<th>Loongana 001</th>
<th>LoV 051</th>
<th>NWA 033</th>
<th>NWA 13400</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>58.5 ± 0.7</td>
<td>58.2 ± 0.8</td>
<td>57.6 ± 0.7</td>
<td>58.4 ± 0.9</td>
<td>58.0 ± 1.0</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.12 ± 0.06</td>
<td>0.10 ± 0.05</td>
<td>0.13 ± 0.06</td>
<td>0.10 ± 0.06</td>
<td>0.10 ± 0.06</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>0.81 ± 0.25</td>
<td>0.59 ± 0.23</td>
<td>0.67 ± 0.24</td>
<td>0.73 ± 0.30</td>
<td>0.71 ± 0.16</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.37 ± 0.11</td>
<td>0.33 ± 0.07</td>
<td>0.32 ± 0.07</td>
<td>0.39 ± 0.09</td>
<td>0.40 ± 0.07</td>
</tr>
<tr>
<td>FeO</td>
<td>6.37 ± 2.59</td>
<td>6.99 ± 1.40</td>
<td>6.69 ± 2.26</td>
<td>5.32 ± 2.04</td>
<td>4.51 ± 2.55</td>
</tr>
<tr>
<td>MnO</td>
<td>0.14 ± 0.08</td>
<td>0.16 ± 0.05</td>
<td>0.16 ± 0.08</td>
<td>0.13 ± 0.06</td>
<td>0.12 ± 0.05</td>
</tr>
<tr>
<td>MgO</td>
<td>34.2 ± 1.8</td>
<td>33.5 ± 0.9</td>
<td>34.2 ± 1.6</td>
<td>34.6 ± 1.4</td>
<td>35.7 ± 1.5</td>
</tr>
<tr>
<td>CaO</td>
<td>0.48 ± 0.07</td>
<td>0.44 ± 0.15</td>
<td>0.57 ± 0.16</td>
<td>0.45 ± 0.12</td>
<td>0.55 ± 0.05</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.01 ± 0.01</td>
<td>0.01 ± 0.01</td>
<td>0.01 ± 0.02</td>
<td>0.02 ± 0.02</td>
<td>0.02 ± 0.02</td>
</tr>
<tr>
<td>NiO</td>
<td>0.02 ± 0.04</td>
<td>0.05 ± 0.05</td>
<td>0.03 ± 0.02</td>
<td>0.04 ± 0.03</td>
<td>0.09 ± 0.08</td>
</tr>
<tr>
<td>Total</td>
<td>101.02</td>
<td>100.37</td>
<td>100.68</td>
<td>100.18</td>
<td>100.20</td>
</tr>
<tr>
<td>Fs</td>
<td>9.4 ± 3.9 (1.3–14.1)</td>
<td>10.4 ± 2.1 (6.9–14.0)</td>
<td>10.2 ± 3.3 (2.7–14.6)</td>
<td>7.9 ± 3.1 (3.1–15.2)</td>
<td>6.6 ± 3.7 (1.2–15.9)</td>
</tr>
<tr>
<td>Wo</td>
<td>0.9 ± 0.1 (0.7–1.2)</td>
<td>0.9 ± 0.3 (0.5–1.5)</td>
<td>1.1 ± 0.3 (0.7–1.9)</td>
<td>0.9 ± 0.2 (0.4–1.3)</td>
<td>1.0 ± 0.1 (0.8–1.2)</td>
</tr>
<tr>
<td>FeO/MnO</td>
<td>46</td>
<td>44</td>
<td>44</td>
<td>41</td>
<td>38</td>
</tr>
<tr>
<td>n</td>
<td>27</td>
<td>24</td>
<td>45</td>
<td>49</td>
<td>20</td>
</tr>
</tbody>
</table>

Fs, Wo: Ferrosilite, wollastonite content in low-Ca pyroxene (mol%); ±: 1 standard deviation; Numbers in parenthesis: range of measured values.

### Table 6
Mean chemical composition of Ca-rich pyroxene in the investigated CL chondrites (EMPA; wt.%).

<table>
<thead>
<tr>
<th></th>
<th>Coolidge</th>
<th>Loongana 001</th>
<th>LoV 051</th>
<th>NWA 033</th>
<th>NWA 13400</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>54.6 ± 1.1</td>
<td>54.2 ± 2.1</td>
<td>53.7 ± 1.2</td>
<td>53.9 ± 0.7</td>
<td>54.3 ± 0.9</td>
</tr>
<tr>
<td>TiO₂</td>
<td>1.09 ± 0.33</td>
<td>0.93 ± 0.48</td>
<td>0.96 ± 0.31</td>
<td>1.07 ± 0.29</td>
<td>1.06 ± 0.38</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>2.75 ± 1.15</td>
<td>3.91 ± 2.40</td>
<td>3.38 ± 1.40</td>
<td>3.33 ± 1.32</td>
<td>2.91 ± 0.89</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.69 ± 0.16</td>
<td>0.84 ± 0.46</td>
<td>0.69 ± 0.22</td>
<td>0.98 ± 0.31</td>
<td>0.59 ± 0.14</td>
</tr>
<tr>
<td>FeO</td>
<td>4.12 ± 1.97</td>
<td>3.40 ± 2.03</td>
<td>3.42 ± 1.80</td>
<td>2.95 ± 1.36</td>
<td>2.64 ± 1.85</td>
</tr>
<tr>
<td>MnO</td>
<td>0.21 ± 0.05</td>
<td>0.22 ± 0.10</td>
<td>0.17 ± 0.04</td>
<td>0.23 ± 0.08</td>
<td>0.16 ± 0.05</td>
</tr>
<tr>
<td>MgO</td>
<td>21.0 ± 2.5</td>
<td>22.2 ± 4.5</td>
<td>20.6 ± 2.2</td>
<td>19.4 ± 1.6</td>
<td>23.1 ± 3.4</td>
</tr>
<tr>
<td>CaO</td>
<td>16.3 ± 4.2</td>
<td>14.7 ± 6.1</td>
<td>17.3 ± 3.6</td>
<td>18.4 ± 2.6</td>
<td>15.2 ± 4.4</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.30 ± 0.14</td>
<td>0.19 ± 0.14</td>
<td>0.29 ± 0.18</td>
<td>0.33 ± 0.23</td>
<td>0.09 ± 0.12</td>
</tr>
<tr>
<td>NiO</td>
<td>0.03 ± 0.04</td>
<td>0.04 ± 0.07</td>
<td>0.03 ± 0.04</td>
<td>0.04 ± 0.04</td>
<td>0.07 ± 0.05</td>
</tr>
<tr>
<td>Total</td>
<td>101.09</td>
<td>100.63</td>
<td>100.54</td>
<td>100.63</td>
<td>100.12</td>
</tr>
<tr>
<td>Fs</td>
<td>6.6 ± 3.2 (1.4–10.8)</td>
<td>5.4 ± 3.1 (1.6–9.8)</td>
<td>5.3 ± 2.9 (1.4–12.0)</td>
<td>4.8 ± 2.2 (0.7–8.6)</td>
<td>4.2 ± 2.9 (0.8–10.2)</td>
</tr>
<tr>
<td>Wo</td>
<td>33.5 ± 8.8 (18.4–43.6)</td>
<td>30.9 ± 13.5 (12.3–43.7)</td>
<td>35.5 ± 7.6 (16.3–45.6)</td>
<td>38.6 ± 5.4 (27.1–49.3)</td>
<td>30.9 ± 9.4 (10.7–44.6)</td>
</tr>
<tr>
<td>FeO/MnO</td>
<td>20</td>
<td>15</td>
<td>20</td>
<td>13</td>
<td>17</td>
</tr>
<tr>
<td>n</td>
<td>10</td>
<td>8</td>
<td>38</td>
<td>28</td>
<td>23</td>
</tr>
</tbody>
</table>

Fs, Wo: Ferrosilite, wollastonite content in Ca-rich pyroxene (mol%); ±: 1 standard deviation; Numbers in parenthesis: range of measured values.
Table 7

<table>
<thead>
<tr>
<th></th>
<th>Coolidge</th>
<th>Loongana 001</th>
<th>LoV 051</th>
<th>NWA 033</th>
<th>NWA 13400</th>
<th>Mean CL</th>
<th>Allende CV,ox,d</th>
<th>NWA 7020 CR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>404 / (610)</td>
<td>414 / (700)</td>
<td>476</td>
<td>455</td>
<td>477</td>
<td>457</td>
<td>470</td>
<td>633</td>
</tr>
<tr>
<td>Median</td>
<td>270</td>
<td>288</td>
<td>403</td>
<td>332</td>
<td>342</td>
<td>341</td>
<td>357</td>
<td>506</td>
</tr>
<tr>
<td>StDev(^2)</td>
<td>405</td>
<td>347</td>
<td>327</td>
<td>349</td>
<td>421</td>
<td>375</td>
<td>366</td>
<td>468</td>
</tr>
<tr>
<td>Skewness</td>
<td>2.60</td>
<td>2.13</td>
<td>2.07</td>
<td>1.69</td>
<td>2.02</td>
<td>2.07</td>
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<td>1.20</td>
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<tr>
<td>Min(^3)</td>
<td>55</td>
<td>75</td>
<td>75</td>
<td>65</td>
<td>50</td>
<td>50</td>
<td>70</td>
<td>65</td>
</tr>
<tr>
<td>Max(^4)</td>
<td>2970</td>
<td>2295</td>
<td>2490</td>
<td>2070</td>
<td>2930</td>
<td>2930</td>
<td>2670</td>
<td>2540</td>
</tr>
<tr>
<td>n(^5)</td>
<td>145</td>
<td>235</td>
<td>397</td>
<td>274</td>
<td>512</td>
<td>1563</td>
<td>421</td>
<td>148</td>
</tr>
</tbody>
</table>

1. Literature data (Kallemeyn and Rubin, 1995).
2. 1 standard deviation.
3. Size of the smallest chondrule.
4. Size of the largest chondrule.
5. Number of analyzed chondrules.

The measured size-frequency distributions of the investigated CL chondrites are shown in Fig. 5a. In Fig. 5b their mean size distribution is compared to the distributions found in the present study for the CV chondrite Allende and the CR chondrite NWA 7020. The mean CL distribution is similar to that of Allende, but the CR distribution differs in having much larger fractions of large (>600 μm) chondrules and correspondingly lower fractions of smaller chondrules.

4.7.2. Refractory inclusions (CAIs and AOAs)

The largest CAIs found in our study are from NWA 13400 and Loongana, with maximum apparent sizes of 3.8 and 2.8 mm, respectively. The occurrence of CAIs in Loongana and Coolidge has been described by Noguchi (1994) and Kallemeyn and Rubin (1995) with maximum sizes of 500 μm in Coolidge. All CAIs in our samples are fine-grained and consist mainly of an intergrowth of spinel, plagioclase, and Ca-rich pyroxene. Similar CAI textures were also described by Noguchi (1994) for the Coolidge meteorite. A complex CAI in Loongana is shown in Fig. 6a, which consists of fine-grained subunits (anorthite, spinel), rimmed by Ca-rich pyroxene. Some FeS is also present in this CAI. Several other CAIs are rich in FeS, probably a result of secondary alteration on the parent asteroid. An example is shown in Fig. 6b from NWA 13400, which consists of anhedral to euhedral spinel grains, embedded in a fine-grained groundmass of spinel, FeS, and plagioclase. This inclusion is partially rimmed by Ca-rich pyroxene. Spinel grains in all CAIs are zoned, with decreasing Mg- and Al-contents and increasing Cr- and Fe-contents from core to rim. This confirms observations by Noguchi (1994) for Coolidge. In a CAI from this meteorite we found an assemblage of spinel + FeS + TiO₂ (probably rutile; Fig. 6c). This CAI is rimmed by an outer layer of diopside and an inner layer of spinel, possibly remnants of a former Wark-Lovering rim. CAIs in the investigated meteorites are frequently accompanied by large grains of Ca-apatite, either on their edges (up to 500 μm) or in their interiors (up to 100 μm; Fig. 6d). These phosphates possibly formed by the release of Ca during alteration of CAIs. This relation was also observed in Coolidge by Noguchi (1994). Moreover, we found a small (7 μm) refractory metal nugget with high concentrations of Os, Ir, Pt, and Ru in a CAI in
Loongana, placed within a region of altered Fe-Ni metal (Fig. 6e).

Small (<a few hundred μm) amoeboid olivine aggregates (AOAs) were found in Coolidge by Noguchi (1994). We found a large AOA with an apparent maximum size of 900 μm in LoV 051 (Fig. 6f), which consists of a porous part rich in olivine and a more compact part that contains small CAI-like units. These units mainly consist of Ca-rich plagioclase (An70) and chromian spinel.

4.7.3. Matrix

The modal amount of matrix is low in the investigated samples (17–21 vol% Table 2). The matrix mainly consists of an assortment of olivine grains with smaller amounts of low-Ca pyroxene and some plagioclase. Similar observations were described by Noguchi (1994) for the Coolidge meteorite. Fe-rich materials fills the pore spaces and probably represent oxidation products of metals and sulfides (Fig. 7a), a conclusion also given in McSween (1977a). The grain size distribution of matrix components is bimodal, where larger grains (~2–30 μm) are embedded in a groundmass of smaller grains with sizes mostly between ~0.2 and ~1 μm (Fig. 7b–f). The majority of larger grains represent fragments of formerly larger objects, probably chondrules, with fracture surfaces and irregular outlines. Beside some angular fragments, the smaller grains are a
mixture of isometric to slightly elongated grains, which may have crystallized during thermal metamorphism. The mean chemical compositions of matrix in the investigated samples, obtained by EMPA, are listed in Table 8, together with literature data for other carbonaceous chondrite groups. To provide a better comparison with our values, the literature data of McSween and Richardson (1977) in this table have been recalculated to totals of 95%.

### 4.8. Bulk meteorite elemental compositions

#### 4.8.1. Major and minor elements

The measured bulk concentrations of major and minor elements in the CL chondrites and the Allende reference sample are listed in Table 9, together with literature data for Coolidge, Loongana, and Allende. The measured values for Allende are well within or close to the range of literature values. Our FeO (30.99 wt.%) and Na₂O (0.32 wt.%) values for Coolidge appear high compared to literature values (24.83–27.04 and 0.24–0.26 wt.%, respectively). However, our values for FeO and Na₂O of the Allende reference sample are in the range of literature data, thus showing no systematic analytical bias. None of the chosen analysis methods of this study was able to measure the bulk S content of the samples and we used SEM-EDX data to obtain a rough estimate of the S concentrations (see Section 3.2.1). These results are also included in Table 9.

Regarding the concentration ratios of the lithophile elements Si, Mg, and Al, CL data plot within the field of the other carbonaceous chondrite groups (Fig. 8a). Compared to Allende, the CL chondrites are slightly depleted in Al and K and strongly depleted in Mn and Na. Hence, they are chemically well discriminated from CV chondrites and the other groups of carbonaceous chondrites (e.g., Fig. 8b).

#### 4.8.2. Trace elements

The measured bulk concentrations of trace elements in the samples and in the Allende reference sample are listed in Table 10, together with literature data for Coolidge, Loongana, and Allende. Our measurements are plotted in Fig. 8c and d and Figs. 9 and 10, together with data for major and minor elements. The measured values for Allende are well within or close to the range of literature values (Table 10). One exception is Te, for which we determined a value of 1.21 ppm, compared to literature values between 1.04 ppm (Braukmüller et al., 2018) and 0.94 ppm (Wang et al., 2014). Thus, our Te values for the CL samples may be systematically too high (16%–29%). As noted above, severe weathering effects were detected for Loongana and NWA 033. For example, in these meteorites the light rare earth elements (REE) were enriched during terrestrial weathering, a process described by, e.g., Pourkhorsandi et al. (2017a). These data and some for the other samples are shown in italics in Table 10 and are neither included in the figures nor will they be included in the discussion.

Only few literature data exist on the chemical composition of Coolidge (Kallemeyn and Rubin, 1995; Weckwerth, 2014; Table 10). Compared to these, our values for Sc and Se are higher and those for V and Hf are somewhat lower (Table 10). Compared to literature data (Kallemeyn and Rubin, 1995), our Loongana value for Se is somewhat higher and those for V and Zn are somewhat lower. Again, since our values for the respective elements in the Allende reference sample are in the range of literature data, there appears to be no systematic analytical bias. The investigated CL chondrites are chemically well discernable from the other carbonaceous chondrite groups on Al₂O₃ vs. Zn (Fig. 8c) and Zr vs. Rb (Fig. 8d) plots, which reveal the characteristic depletion of volatile elements such as Rb and Zn.

### Table 8

Bulk composition of Matrix in the investigated CL chondrites (EMPA; wt.%). Literature values for other carbonaceous groups are given for comparison. Clg: Coolidge; Loo: Loongana; LoV: LoV051; 033: NWA 033; 13400: NWA 13400.

<table>
<thead>
<tr>
<th>Element</th>
<th>Clg</th>
<th>Loo</th>
<th>LoV</th>
<th>033</th>
<th>13400</th>
<th>Mean CL</th>
<th>Vigarano*</th>
<th>Allende*</th>
<th>Karoonda*</th>
<th>Renazzo*</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>30.8</td>
<td>27.7</td>
<td>30.9</td>
<td>29.5</td>
<td>32.8</td>
<td>30.3</td>
<td>31.2</td>
<td>30.1</td>
<td>34.1</td>
<td>36.6</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.10</td>
<td>0.13</td>
<td>0.13</td>
<td>0.11</td>
<td>0.18</td>
<td>0.13</td>
<td>0.10</td>
<td>0.10</td>
<td>0.55</td>
<td>0.08</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>2.01</td>
<td>2.18</td>
<td>1.56</td>
<td>2.04</td>
<td>1.41</td>
<td>1.84</td>
<td>4.70</td>
<td>2.48</td>
<td>7.63</td>
<td>3.10</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.79</td>
<td>0.81</td>
<td>0.44</td>
<td>0.61</td>
<td>0.89</td>
<td>0.71</td>
<td>0.44</td>
<td>0.41</td>
<td>0.40</td>
<td>0.41</td>
</tr>
<tr>
<td>FeO</td>
<td>31.5</td>
<td>34.5</td>
<td>30.3</td>
<td>32.3</td>
<td>25.2</td>
<td>30.8</td>
<td>35.9</td>
<td>34.3</td>
<td>28.1</td>
<td>28.3</td>
</tr>
<tr>
<td>MnO</td>
<td>0.14</td>
<td>0.15</td>
<td>0.16</td>
<td>0.15</td>
<td>0.20</td>
<td>0.16</td>
<td>0.15</td>
<td>0.23</td>
<td>0.14</td>
<td>0.38</td>
</tr>
<tr>
<td>MgO</td>
<td>28.3</td>
<td>25.6</td>
<td>28.9</td>
<td>27.5</td>
<td>31.3</td>
<td>28.3</td>
<td>18.8</td>
<td>21.7</td>
<td>18.7</td>
<td>18.4</td>
</tr>
<tr>
<td>CaO</td>
<td>0.97</td>
<td>1.11</td>
<td>1.11</td>
<td>1.23</td>
<td>0.98</td>
<td>1.18</td>
<td>1.68</td>
<td>2.55</td>
<td>3.98</td>
<td>1.01</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.25</td>
<td>0.29</td>
<td>0.36</td>
<td>0.53</td>
<td>0.09</td>
<td>0.30</td>
<td>0.54</td>
<td>0.24</td>
<td>0.22</td>
<td>1.35</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
<td>0.04</td>
<td>0.03</td>
<td>0.03</td>
<td>0.06</td>
<td>0.01</td>
<td>0.13</td>
<td>0.19</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.15</td>
<td>0.19</td>
<td>0.13</td>
<td>0.16</td>
<td>0.24</td>
<td>0.17</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>Ni</td>
<td>1.01</td>
<td>1.56</td>
<td>1.04</td>
<td>1.26</td>
<td>0.87</td>
<td>1.15</td>
<td>1.20</td>
<td>1.55</td>
<td>0.22</td>
<td>1.36</td>
</tr>
<tr>
<td>S</td>
<td>0.15</td>
<td>0.09</td>
<td>0.35</td>
<td>0.08</td>
<td>0.04</td>
<td>0.14</td>
<td>0.22</td>
<td>1.22</td>
<td>0.09</td>
<td>3.72</td>
</tr>
<tr>
<td>Total</td>
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<td>94.84</td>
<td>95.41</td>
<td>95.51</td>
<td>94.23</td>
<td>95.24</td>
<td>95.00</td>
<td>95.00</td>
<td>95.00</td>
<td>95.00</td>
</tr>
<tr>
<td>n</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>12</td>
<td>10</td>
<td>20</td>
<td>20</td>
<td>20</td>
<td>20</td>
<td>20</td>
</tr>
</tbody>
</table>

nd: not determined.

n: number of analyzed areas.

1 defocused beam (30 μm)

2 CV subgroups: CV_red = reduced; CV_oxA = oxidized Allende-type (Weisberg et al., 1997).

* McSween and Richardson (1977); recalculated to 95.00 wt.% for comparison.
4.9. Bulk meteorite isotope compositions

4.9.1. Oxygen isotopes

The measured values for \( \delta^{17} \text{O}, \delta^{18} \text{O}, \) and \( \Delta^{17} \text{O} \) of the investigated CL chondrites and Allende are listed in Table 11 and plotted in Fig. 11. The compositional fields for CR, CV3, CK, and CO3 chondrites are shown for comparison. All CL samples plot along the Carbonaceous Chondrite Anhydrous Mineral (CCAM) line (Clayton and Mayeda, 1999) with \( \Delta^{17} \text{O} \) values between \(-3.96\) and \(+5.47\)%e. The CL data plot either directly on this line (NWA 033, NWA 13400) or slightly off it in the direction of the Primitive Chondrule Minerals (PCM) line (Ushikubo et al., 2011). Three of the CL samples (Loongana, NWA 033, NWA 13400) plot within the range of CV3 and CK chondrites, while Coolidge and LoV 051 have considerably lower \( ^{17} \text{O}/^{16} \text{O} \) and \( ^{18} \text{O}/^{16} \text{O} \) ratios.

4.9.2. Titanium and chromium isotopes

The Ti and Cr isotopic data are provided in Table 12 and visualized in Fig. 12. In Fig. 12 a the \( \epsilon^{54} \text{Cr} \) data are plotted versus the corresponding \( \Delta^{17} \text{O} \) values and in Fig. 12 b the \( \epsilon^{54} \text{Cr} \) values are plotted versus the \( \epsilon^{50} \text{Ti} \) data. In such diagrams the dichotomy between non-carbonaceous and carbonaceous chondrites is strikingly evident (Warren, 2011; Kleine et al., 2020). Although the \( \epsilon^{54} \text{Cr} \) values of the five CL chondrites are similar to the mean values of CO and CV chondrites (Fig. 12 a), the CL
chondrites plot in a distinctly different region of the $\varepsilon^{54}\text{Cr}$-$\varepsilon^{50}\text{Ti}$ diagram, where they do not overlap with the compositions of CO and CV chondrites (Fig. 12b). Instead, in terms of $\varepsilon^{50}\text{Ti}$ the CL chondrites are more similar to CM and CR chondrites. All CL chondrites have indistinguishable $\varepsilon^{54}\text{Cr}$ and $\varepsilon^{50}\text{Ti}$ values, and so the mean Ti and Cr isotope composition of the CL group (Table 12) can be precisely defined from these data. This in turn makes it possible to use these data for evaluating the genetic relationship of CL chondrites relative to other carbonaceous chondrite groups.

5. DISCUSSION

5.1. Characteristics of CL chondrites and differences to other chondrite groups

The investigated samples share many textural, chemical, and isotopic properties. Macroscopically they remotely resemble CV$_{\text{red}}$ and CR chondrites, but there are many differences to these and other carbonaceous chondrite groups, as discussed in the following.

5.1.1. Chondrules

Most chondrules are PO and POP varieties of Type-I, rich in Fe-Ni metal. Many small chondrules show a remarkably irregular outer shape with lobate outlines and protuberances (Fig. 3a) or even have an amoeboidal shape (Fig. 3b and c). Many of the small chondrules are concentrated in layers around larger chondrules (Fig. 3d). However, such textures are not unique to these samples, but were occasionally observed by us in the CV$_{\text{red}}$ chondrite Vigarano and the CR chondrite NWA 7020, as well. Lobate chondrules also exist in CO chondrites and are explained as having formed from reheated chondrule fragments, i.e., by rounding of fracture surfaces by minor melting (Rubin and Wasson, 2005). Based on the refractory element diversity of different lobes in a given chondrule, Jacquet (2021) concluded that lobate chondrules formed by the collision of preexisting droplets. A formation of lobate chondrules in
Table 10
Bulk chemical composition of the investigated samples (trace elements; ppm). Numbers in italics indicate values that show evidence for modification by terrestrial weathering. Sample abbreviations: see Table 8.

<table>
<thead>
<tr>
<th></th>
<th>Lab¹</th>
<th>RSD²(%)</th>
<th>Clg³</th>
<th>Loo⁴</th>
<th>LoV</th>
<th>033</th>
<th>13400</th>
<th>Mean CL⁵</th>
<th>Allende</th>
<th>Allende (Literature)⁶</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li</td>
<td>B</td>
<td>1.7</td>
<td>2.10</td>
<td>2.13</td>
<td>2.07</td>
<td>2.10</td>
<td>1.97</td>
<td>1.69</td>
<td></td>
<td>1.81³; 1.64²</td>
</tr>
<tr>
<td>Sc</td>
<td>D</td>
<td>4.9</td>
<td>15.5</td>
<td>10.5</td>
<td>15.3</td>
<td>10.6</td>
<td>11.5</td>
<td>12.7</td>
<td>11.8</td>
<td>11.3²; 10.8³; 10.7⁴</td>
</tr>
<tr>
<td>V</td>
<td>B</td>
<td>2.2</td>
<td>85.6</td>
<td>73.3</td>
<td>85.5</td>
<td>81.4</td>
<td>81.6</td>
<td>95.9</td>
<td>90.2*</td>
<td>92.6; 95.2⁵ 91³ 91⁶</td>
</tr>
<tr>
<td>Zn</td>
<td>A</td>
<td>&lt; 1</td>
<td>56.7</td>
<td>33.7</td>
<td>32.5</td>
<td>43.5</td>
<td>51.6</td>
<td>43.6</td>
<td>102</td>
<td>106; 123 119³ 118⁴</td>
</tr>
<tr>
<td>Sc</td>
<td>D</td>
<td>32</td>
<td>5.05</td>
<td>5.78</td>
<td>2.62</td>
<td>5.18</td>
<td>3.04</td>
<td>4.33</td>
<td>9.57</td>
<td>9.21³; 8.25⁵</td>
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<tr>
<td>Rb</td>
<td>B</td>
<td>2.3</td>
<td>0.29</td>
<td>0.92</td>
<td>0.42</td>
<td>0.50</td>
<td>0.18</td>
<td>0.46</td>
<td>1.21</td>
<td>1.32 1.15         1.18³ 1.17²</td>
</tr>
<tr>
<td>Sr</td>
<td>B</td>
<td>0.6</td>
<td>14.8</td>
<td>15.3</td>
<td>18.8</td>
<td>24.2</td>
<td>15.5</td>
<td>16.1</td>
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<tr>
<td>Y</td>
<td>B</td>
<td>1.2</td>
<td>2.66</td>
<td>3.75</td>
<td>2.81</td>
<td>3.77</td>
<td>2.48</td>
<td>3.09</td>
<td>2.60</td>
<td>2.63 2.77</td>
</tr>
<tr>
<td>Zr</td>
<td>B</td>
<td>0.7</td>
<td>6.82</td>
<td>8.89</td>
<td>7.40</td>
<td>9.15</td>
<td>6.20</td>
<td>7.69</td>
<td>6.60</td>
<td>6.77 6.45           6.61³ 7.1³</td>
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<tr>
<td>Nb</td>
<td>B</td>
<td>2.4</td>
<td>0.519</td>
<td>0.620</td>
<td>0.662</td>
<td>0.574</td>
<td>0.535</td>
<td>0.582</td>
<td>0.531</td>
<td>0.569 0.503         0.57³ 0.58⁵</td>
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CL chondrites by droplet collision would fit well to the observed chondrule textures (e.g., Fig. 3d). A distinctive feature of the CL samples is their high chondrule abundance (67 to 79 vol%; Table 2). Kallemeyn and Rubin (1995) found nearly the same chondrule abundance in Loongana (~70% vs. 67% in this study), but a lower abundance in Coolidge (60–65% vs. 79% in this study), possibly reflecting sample heterogeneity. McSween (1977a) found 67 vol% chondrules in Coolidge. For comparison, chondrule concentrations in CV and CR chondrites are 45% and 50–60%, respectively (Krot et al., 2014), i.e., considerably lower than in CL chondrites.

5.1.2. Apparent chondrule sizes

The observed range of mean apparent chondrule sizes in the CL samples is restricted to values between 404 and 477 μm (Table 7). To verify our method, we also measured the apparent (2D) chondrule sizes in the CR2 chondrite NWA 7020 and the CV OxA chondrite Allende (Table 7; Fig. 5b). It is found that the values of mean chondrule sizes in CLs overlap with those in the CV chondrite Allende (470 μm; Table 7). Other statistical parameters of the CL chondrites size-frequency distributions (median size, standard deviation, and skewness) are also similar to those displayed by Allende (Table 7), hinting at a remote relationship to CV chondrites. The CL mean sizes are distinctly smaller than that of the CR chondrite NWA 7020, which shows a mean size of 633 μm and a much more symmetric distribution (skewness of only 1.20, compared to 1.69–2.60 in CLs; Table 7).

Our mean value for the CR2 chondrite (633 μm) is in the range of published values for five other CR chondrites (490–770 μm; Kallemeyn and Rubin, 1995) larger than those found by us (404 μm and 414 μm). The likely explanation is that we measured a large number of chondrules (145 and 235; Table 7) on BSE photomosaics, whereas the literature values were obtained from only 23–52 chondrules (Kallemeyn et al., 1994) using optical microscopy. This could have led to an overrepresentation of large chondrules in their statistics.

From the presented results it can be concluded that the chondrule-size-frequency distributions and mean chondrule

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<tr>
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1 Analytical method and laboratory: A: XRF, Institut für Geologie und Mineralogie, University of Cologne; B: ICP-MS, Institut für Mineralogie, University of Münster, Germany; D: LA-ICP-MS, Institut für Mineralogie, University of Münster.
2 Reproducibility of the measured values (in %, RSD).
3 Literature data (Kallemeyn and Rubin, 1995; Weckwerth, 2014) in parenthesis; nd: not determined.
4 Literature data (Kallemeyn and Rubin, 1995) in parenthesis.
5 Numbers in italics (individual meteorite data) have been excluded from mean value calculation.
6 Literature data for Allende. Averaged values are shown in Figs. 9 and 10, * XRF/ICP-MS average; † XRF/ICP-MS average; ‡ XRF/ICP-MS average.
7 Literature data (Kallemeyn and Rubin, 1995) in parenthesis.
8 Literature data (Kallemeyn and Rubin, 1995) in parenthesis. The observed range of mean apparent chondrule sizes in the CL samples is restricted to values between 404 and 477 μm (Table 7). To verify our method, we also measured the apparent (2D) chondrule sizes in the CR2 chondrite NWA 7020 and the CV chondrite Allende (Table 7; Fig. 5b). It is found that the values of mean chondrule sizes in CLs overlap with those in the CV chondrite Allende (470 μm; Table 7). Other statistical parameters of the CL chondrites size-frequency distributions (median size, standard deviation, and skewness) are also similar to those displayed by Allende (Table 7), hinting at a remote relationship to CV chondrites. The CL mean sizes are distinctly smaller than that of the CR chondrite NWA 7020, which shows a mean size of 633 μm and a much more symmetric distribution (skewness of only 1.20, compared to 1.69–2.60 in CLs; Table 7).

Literature values of mean apparent chondrule sizes for Coolidge and Loongana (610 μm and 700 μm; Kallemeyn and Rubin, 1995) are larger than those found by us (404 μm and 414 μm). The likely explanation is that we measured a large number of chondrules (145 and 235; Table 7) on BSE photomosaics, whereas the literature values were obtained from only 23–52 chondrules (Kallemeyn et al., 1994) using optical microscopy. This could have led to an overrepresentation of large chondrules in their statistics.

Our mean value for the CR2 chondrite (633 μm) is in the range of published values for five other CR chondrites (490–770 μm) and close to the corresponding average value (658 μm; Kallemeyn et al., 1994). In case of Allende a verification of our mean value (470 μm) is more complicated since literature data vary considerably, with mean values of 915 μm (Teitler et al., 2010), 765 μm (Gattacceca et al., 2020), and 310 μm (Simon et al., 2018). Sample inhomogeneity may be partly responsible for these variations. Furthermore, the different methods of investigation certainly contribute to these differences, i.e., chondrule separation (true chondrule sizes; Teitler et al., 2010), optical microscopy (apparent chondrule sizes; Gattacceca et al., 2020), and scanning electron microscopy (apparent chondrule sizes; x-ray compositional images, Simon et al., 2018; BSE images, this work). Furthermore, the classification of small subrounded particle cut faces either as chondrules or fragments is a source of subjective bias.
sizes in CL chondrites are similar to each other and to those in CV chondrites, but dissimilar to those in CR chondrites (Fig. 5; Table 7).

5.1.3. Mean olivine compositions

Similar mean compositions of olivine (Fa 12.5–14.7 mol %; Table 4) are a further distinctive feature of CL chondrites. However, this property is of secondary origin, established by thermal metamorphism on the parent body. This cannot be a criterion for recognizing unequilibrated CL samples that may be identified in the future (see Section 5.3.3). On the other hand, this characteristic range of mean Fa values discerns the CL samples from metamorphosed CV and CR meteorites, since mean Fa values in the latter are much higher. For example, olivine in the CV4 chondrite NWA 8418 (Mallozzi et al., 2018), originally classified as C3 (MBDB), shows a mean Fa value of ~40 mol%, with a pronounced peak at ~38 mol%, far away from the Fa values in CL samples. The CR6 chondrites NWA 6921, NWA 7317, and NWA 11561 show mean Fa values between 29 and 38 mol% (MBDB), which is also much higher than in CL chondrites. Due to these differences it is obvious that CL chondrites do not represent thermally altered CV and CR chondrites.

5.1.4. Fe-Ni metal abundance

Another distinctive feature of CL compared to CV chondrites is the high amount of Fe-Ni metal, which influences their magnetic properties. The magnetic susceptibility of LoV 051, the least weathered sample, is \( \log_\gamma = 5.24 \) (where \( \gamma \) is in \( 10^{-9} \text{ m}^3 \text{ kg}^{-1} \)), which is similar to the mean value for CR chondrites of \( \log_\gamma = 5.04 \pm 0.12 \) (Table 3; Rocchette et al., 2008). The value for the saturation magnetization \( M_S \) (32.3 Am\(^2\)/kg) translates into a kamacite content of 14.4 wt.% for LoV 051. The hysteresis properties of the investigated samples are similar to those of Renazzo CR chondrite. They are markedly different from those of CV chondrites (Weisberg et al., 1997), including CV\(_{\text{red}}\) that

Fig. 9. CI-normalized (Lodders et al, 2009) lithophile element concentrations in CL chondrites and other carbonaceous chondrite groups. Elements are arranged in the order of increasing volatility (Wood et al. 2019). (a) Data for CL chondrites (colored curves, circles) and Allende (black curve, squares), together with Allende literature data (dashed curve; see Tables 9 and 10). The volatile lithophile elements Mn, Na, K, Rb, and Cs are moderately to strongly depleted compared to Allende. (b) Mean concentrations in the CL chondrites (red solid curve, filled circles) and other carbonaceous chondrite groups (CM, CO, CV, CR, CK4; literature values; black curves). The CL-trend of volatile element depletion and enrichment of other elements is evident. Literature data (mean values, where possible): Knab (1981); Kallemeyn and Wasson (1981); Jarosewich et al. (1987); Kallemeyn et al. (1991, 1994); Wolf and Palme (2001); Jarosewich (2006); Greenwood et al. (2010); McCoy-West et al. (2017); Braukmüller et al. (2018).
show the highest Fe-Ni metal abundances. Indeed, CV red meteorites contain much less metal than LoV 051, i.e., 2.73 ± 1.12 vol% (average ± s.d., n = 9; Bonal et al., 2020), translating into 6.8 ± 2.8 wt.%.

5.1.5. CAI abundances

The CAI abundances appear to be another characteristic property of the CL chondrites, although our data base is limited. The modal abundance of CAIs is highly variable (from 0.1 vol% for Coolidge to 8 vol% for Loongana; Table 2), probably due to the small section sizes (52–361 mm²; Table 1). It was shown by Hezel et al. (2008) that section sizes <1000 mm² do not give representative results, as CAI abundances follow a Poisson distribution in thin sections. A clear hint on this is the observation that the sample with the apparently highest CAI abundance (Loongana) shows the lowest bulk Al₂O₃ concentration (Table 9). Taking all measurements together (655 mm²), the average CAI abundance is 1.4 vol%. Hence, our best guess for the CAI concentration in these samples is 1.4 vol%, which is much higher than in CR chondrites (0.12 vol%) and distinctly lower than in CV chondrites (3 vol% or 4 vol%; Hezel et al., 2008; Ebel et al., 2016). Our estimate is comparable to that given in Kallemeyn and Rubin (1995) for Coolidge and Loongana (1–2 vol%).
5.1.6. Matrix

Complementary to the high chondrule concentration, the low amount of matrix (17–21%; Table 2) is another important distinctive feature of CL chondrites. The mean modal abundances of matrix in CV and CR chondrites are much higher, i.e., 52 vol% and 40 vol% in CVox and CVred, respectively, (Gattacceca et al., 2020) and 35 vol% in CR chondrites (Schrader et al., 2011). Our value of 18% for Coolidge is nearly identical to the value (17.6%) given by McSween (1977a). The value for matrix in Coolidge as obtained by Kallemeyn and Rubin (1995) is higher (30%), but may reflect sample inhomogeneity. Their value for matrix abundance in Loongana is 20%, identical to the value found in the present study (Table 2).

It was found by Noguchi (1994) that Coolidge matrix is depleted in Na, compared to the reduced CV chondrite Efremovka. This can be confirmed by our measurements, since we found a Na2O concentration of 0.25 wt.% in Coolidge matrix, which is only half the concentration found in Vigarano Matrix (0.54 wt.%; McSween and Richardson, 1977; Table 8). However, the Na2O concentration in CL matrix varies considerably between 0.09 wt.% (NWA 13400) and 0.53 wt.% (NWA 033), with a mean value of 0.30 wt.% (Table 8). The mean concentration of Al2O3 in CL chondrite matrix (1.84 wt.%) is considerably lower than in the matrices of CV, CR and CK chondrites (2.48–7.63 wt.%; Table 8). Another striking difference is the enrichment of Cr2O3 (mean value 0.71 wt.%) in the CL chondrite matrix, compared to 0.40–0.44 wt.% in CV, CK, and CR chondrites (Table 8). This characteristic was already mentioned by McSween and Richardson (1977) for the Coolidge chondrite. The clearest difference to matrix compositions of CV, CR, and CK chondrites are the extraordinarily high MgO concentrations in the CL matrices (mean 28.3 wt.%), compared to mean values of 18.4–21.7 wt.% in the other groups (Table 8). Since olivine and pyroxene are the main carriers of Mg, we have to conclude that these minerals occur in higher abundances in CL matrix than in CV, CK, and CR matrix. This may be a primary characteristic of CL matrix or be the result of matrix modification during thermal metamorphism.

The bimodal size distribution of matrix silicates (see Fig. 7) may indicate the growth of small secondary olivine and pyroxene, by which an enrichment of Mg in bulk matrix may be partly explained. Furthermore, S is considerably depleted in the CL matrix (mean value 0.14 wt.%),
compared to CR chondrites and the CV subgroups. Only CK chondrites show a lower mean concentration (0.09 wt.%). However, CL3.9/CL4 and CK4 chondrites were affected by thermal metamorphism on their parent asteroids, by which their fine-grained matrix may have undergone considerable chemical modifications.

5.1.7. Bulk chemical compositions

The bulk chemical compositions of the five investigated CL samples prove that they belong to the class of carbonaceous chondrites. This is shown in Fig. 8a, where the mean bulk Si/Mg versus Al/Mg element ratios of the CL group are plotted (red filled circle). The CL data plot among the mean data for the other major carbonaceous chondrite groups (Hutchison, 2004; his table 2.1), close to the value for the CK group. Ordinary and R chondrites plot close to each other, but in a distinctly different field in the diagram. The mean values for both enstatite chondrite groups (EH and EL) are also located far away from the field for carbonaceous chondrites. The most characteristic chemical difference between CL chondrites and all other carbonaceous chondrite groups (besides CB and CH) is their strong depletion of volatile lithophile and chalcophile elements. The differences to the CV chondrite Allende can be seen in Tables 9 and 10 and Figs. 9 and 10. This distinctive feature was previously noticed for Coolidge and Loongana (e.g. Kalleney and Wasson, 1982; Kalleney and Rubin, 1995; Weckwerth, 2014). The CL chondrites are well resolved from other carbonaceous chondrite groups on, for example Al₂O₃ vs. MnO (Fig. 8b), Al₂O₃ vs. Zn (Fig. 8c), and Zr vs. Rb (Fig. 8d), since volatile elements like Mn, Zn, and Rb are moderately to strongly depleted.

In Fig. 9a the CI-normalized concentrations (Lodders et al., 2009) of all measured lithophile elements in the CL samples and the Allende reference sample are shown (see Table 10), together with Allende literature data (dotted line). Elements in Figs. 9 and 10 are arranged in the order of increasing volatility (Wood et al., 2019). While elements from Zr (most refractory) to ~Eu in the CL chondrites and Allende exhibit similar CI-normalized abundances, elements from V to Cs are increasingly depleted. This trend steepens for elements more volatile than Cr (Mn, Na, K, Rb, and Cs), with much stronger depletions in the CLs compared to Allende. This can also be seen in the mean CL values in Fig. 9b, when compared to mean values for the other carbonaceous chondrite groups (CM, CO, CV, CR, CK4). The CL chondrites are characterized by strong enrichments of refractory and the strongest depletions of volatile elements among all carbonaceous chondrite groups.

Fig. 10 shows the CI-normalized concentrations of siderophile and chalcophile elements in CL chondrites. In Fig. 10a the values for individual CL chondrites are given, again in comparison to Allende values (measurements and literature data). In Fig. 10b the mean CL values are given, compared to the mean values of the other main carbonaceous chondrite groups. For the siderophile elements Mo, Ni, Co, Fe, and P all samples show a similar pattern, which is also similar to that of the Allende chondrite (Fig. 10a). The mean CL values (Fig. 10b) are in the range of those in the other groups, with the highest concentration of Mo, the most refractory element in this plot. The chalcophile element concentrations in the CL chondrites show some scatter, but are consistently lower than in Allende (Fig. 10a). The mean CL values for chalcophile elements are plotted in Fig. 10b, compared to mean values for the other carbonaceous chondrite groups. The CL mean values indicate the strongest depletions of these elements compared to all other groups, which is in line with the strong depletions of volatile lithophile elements (Fig. 9). Recently, Braukmüller et al. (2018) have shown that concentrations of volatile elements with 50% condensation temperatures below 800 K (from Zn to Ti in Fig. 10) systematically differ between the various groups, but are depleted to about the same extent in a given group (Fig. 10b). Therefore, they termed them “plateau volatile elements”. The CL chondrites investigated in this study deviate somewhat from a plateau-like pattern, since the Zn, Pb, and Ti concentrations appear low compared to the Se and Te concentrations (Fig. 10b). However, Braukmüller et al. (2018) also showed that in the ungrouped carbonaceous chondrite EET 96026 the relative concentrations of the plateau volatile elements were strongly modified during thermal metamorphism. As such, the deviations for the CL chondrites observed in this study may also reflect some element fractionations during thermal metamorphism. This issue will be further discussed below (Section 5.2).

5.1.8. Bulk isotopic compositions

Fig. 11 shows that the O isotopic compositions of CL chondrites (Table 11) do not define a tight cluster, but exhibit a range of isotopic compositions along the CCAM line (Δ17O = 3.96 to −5.47‰). This is also typical of the CV and CK groups and may be indicative of some form of relationship between these groups. Two samples (Coolidge, LoV 051) fall outside the compositional fields of CV and CK chondrites and have considerably lower 17O/18O and 18O/16O ratios. However, Gattacceca et al. (2020) presented O isotope data of two Antarctic CVred chondrites, which overlap with our values for Coolidge. On the other hand, the O isotopic compositions of CL chondrites are clearly distinct from CR chondrites, ruling out any genetic link between these two chondrite groups.

The investigated CL samples have a mean ε²⁴Cr value of 0.72 ± 0.08 (Table 12), which is slightly lower than the ε²⁴Cr value of CV chondrites (mean ε²⁴Cr = 0.87 ± 0.07; Burkhardt et al., 2019) and can clearly be distinguished from the mean ε²⁴Cr values of CM and CR chondrites (Fig. 12; Trinquier et al., 2007; Qin et al., 2010). The only known carbonaceous chondrite group with similar ε²⁴Cr are the CO chondrites. By contrast, all CL samples exhibit significantly lower ε⁵⁰Ti values than typically observed for CV, CO, and CK groups (e.g. Trinquier et al., 2009; Zhang et al., 2012; Burkhardt et al., 2019), and their weighted mean ε⁵⁰Ti of 2.60 ± 0.22 (Table 12) is also significantly lower than the ε⁵⁰Ti of 3.21 ± 0.11 measured for bulk Allende (CV3) in the present study. As such, the Ti isotopic composition of CL chondrites is more similar to that of CM and CR chondrites (Fig. 12b). Thus, on a plot of ε⁵⁰Ti vs ε²⁴Cr the CL chondrites define a distinct cluster with a unique position, making this plot particularly useful as a means of demonstrating that the CL chondrites are iso-
topically distinct from the other carbonaceous chondrite groups.

CAIs are highly enriched in Ti and are characterized by large $^{50}\text{Ti}$ excesses (e.g., $^{50}\text{Ti} \approx 9$; Davis et al., 2018; Render et al., 2019), and, as such, exert a strong control on the Ti concentration and Ti isotopic composition of bulk carbonaceous chondrites (e.g., Bürkhardt et al., 2019). For instance, CAI-free CI chondrules display the lowest Ti concentrations and $^{50}\text{Ti}$ excesses among the carbonaceous chondrites, whereas the CAI-rich CV, CO, and CK chondrites have the highest Ti concentrations and exhibit large $^{50}\text{Ti}$ excesses (e.g. Trinquier et al., 2009). As such, the less-pronounced $^{50}\text{Ti}$ excesses of the CL relative to CV, CO, and CK chondrites suggest that CL chondrites are characterized by significantly lower CAI abundances compared to the other groups of carbonaceous chondrites. Note that given the large amount of sample material used for analyses, it is very unlikely that the less-pronounced $^{50}\text{Ti}$ excesses of the CL chondrites of this study reflect a sampling bias, that is, an unrepresentative lower CAI abundance in the selected sample aliquot. Instead, the lower $^{50}\text{Ti}$ of CL compared to CV chondrites is fully in line with petrographic observations indicating that CL chondrites contain only about 1.4 vol% of CAIs (see Sections 4.7.2 and 5.1.5), in contrast to $\sim 3$ vol% in CV chondrites (Hezel et al., 2008). Surprisingly though, CL chondrites have essentially the same Ti content as the CAI-rich CV chondrules, indicating that CL chondrites incorporated a Ti-rich component which, unlike CAIs, is not characterized by large $^{50}\text{Ti}$ excesses. The origin and nature of this component is difficult to assess, but we note that Na-Al-rich chondrules from ordinary chondrites contain a refractory component which is chemically CAI-like, yet is characterized by $^{50}\text{Ti}$ deficits and not excesses (Ebert et al., 2018).

Incorporation of such material into CL chondrites may, therefore, account for their elevated Ti content but lack of more pronounced $^{50}\text{Ti}$ excesses. Contrary to Ti, CAIs are depleted in Cr relative to the major chondrite components (e.g., chondrules and matrix), and so varying CAI abundances have a negligible effect on the Cr isotopic composition of bulk chondrites. Instead, the $^{54}\text{Cr}$ variations among carbonaceous chondrites predominantly reflect variable amounts of chondrules and matrix. For instance, Hellmann et al. (2020) have shown that among the major carbonaceous chondrite groups (except CR) $^{54}\text{Cr}$ is correlated with the mass fraction of matrix in each group (Fig. 13a). These authors have also shown that bulk carbonaceous chondrites plot along a mixing line between the mean composition of CV, CO, and CM chondrules and bulk CI chondrites, indicating that the $^{54}\text{Cr}$ variability reflects different mixtures of $^{54}\text{Cr}$-rich and CI-like matrix and $^{54}\text{Cr}$-poorer chondrules. The CR chondrites plot off this mixing line, because their chondrules have a distinct $^{54}\text{Cr}$ composition, either because they formed from different precursor material and/or because they formed later (Hellmann et al., 2020). Assessing whether the CL chondrites plot on the chondrule-matrix mixing line defined by the other carbonaceous chondrite groups requires conversion of their matrix volume fraction (~19 vol% on average; Table 2) to matrix mass fraction.

Following the approach of Hellmann et al. (2020) this results in an average matrix fraction of 12 wt.% for the CL chondrites. Using this value together with the mean $^{54}\text{Cr}$ of ~0.7 reveals that CL chondrites plot exactly on the mixing line between mean CV, CO, and CM chondrules and bulk CI chondrites (Fig. 13a). On the one hand this indicates that the $^{54}\text{Cr}$ values of CL chondrules are similar to those of the CV, CO, and CM chondrules and on the other hand that the low $^{54}\text{Cr}$ values of bulk CL chondrites reflects their low matrix fraction.

5.2. Volatile depletions: nebular fractionation or thermal metamorphism on the parent body?

The bulk concentrations of volatile elements systemati- cally co-vary among the distinct carbonaceous chondrite groups (CI, CM, CV, CO, CR) and, like the bulk $^{54}\text{Cr}$ values, are positively correlated with the group-specific matrix abundances (Hellmann et al., 2020). This is illustrated in Fig. 13b–d, where the matrix mass fractions of the carbonaceous chondrite groups are plotted against their average concentrations of the volatile elements Zn, Se, and Te. The CL group, together with the other carbonaceous chondrite groups, plot on these correlation lines, corroborating earlier conclusions that the variable volatile depletions among the carbonaceous chondrites predominantly reflect variable matrix abundances (e.g., Alexander, 2019; Hellmann et al., 2020). A corollary of this observation is that the strong volatile depletion of CL chondrites reflects their very low matrix abundance.

Since the investigated CL chondrites were heated on their parent asteroid, we also have to consider the possibility of volatile loss by this process. Such effects were investigated for Allende (CV$_{\alpha\alpha\alpha}$; Wulf et al., 1995) and for Murchison (CM2; Braukmüller et al., 2018) with heating experiments under oxidizing and reducing conditions. The depletion of the volatile lithophile elements Mn, Na, K, Rb, and Cs is one of the most distinctive features of CL chondrites (Figs. 8 and 9). However, Braukmüller et al. (2018) did not observe considerable depletions of these elements during their heating experiments (Mn was not considered). The behavior of Mn was investigated by Wulf et al. (1995) who found that Mn does not show any depletions, even not in the experiments performed at 1300 °C, neither under oxidizing nor reducing conditions. Considerable depletions of chalcophile elements like Zn, Te, Pb, and TI were found by Wulf et al. (1995) and Braukmüller et al. (2018) under reducing conditions. However, the experimentally produced depletion of Pb is higher than for Zn (Braukmüller et al., 2018), whereas these two elements are depleted in CL chondrites by about the same magnitude (Fig. 10). From these observations we conclude that the consistently low bulk concentrations of volatile lithophile elements (Mn, Na, K, Rb, and Cs) and volatile chalcophile elements (Zn, Se, Te, Pb and TI) in CL chondrites are a primary feature and do not result from thermal metamorphism on the parent body. However, it is possible that the latter process affected the chalcophile element concentrations in the CL samples to some extent. In particular, the relative abundances of the plateau volatile elements...
(Zn, Se, Te, Pb, Tl; Braukmüller et al., 2018) appear to have been modified by this process, because the ratios of these elements vary considerably among individual CL samples (Fig. 10a), possibly indicating local redistribution on the parent body.

5.3. Proposal for the new Loongana (CL) group of carbonaceous chondrites

Based on their common characteristics we conclude that the five investigated meteorites can be combined together to form a new group of carbonaceous chondrites. The foundation for this was given by Kallemeyn and Rubin (1995), who defined the Coolidge-Loongana grouplet.

5.3.1. Proposed naming as the CL group of carbonaceous chondrites

Among the investigated samples only Coolidge was given an individual meteorite name; the others are named by serial numbers (Table 1). Hence, the first alphabetic character of Coolidge would be the best candidate to name this group, leading to the CC group of carbonaceous chondrites, which is precluded by the already existing CC meteorite class (CC for carbonaceous chondrites). Since Loongana 001 was the first meteorite for which a close relation to Coolidge was described (Kallemeyn and Rubin, 1995), we propose to use its first alphabetic character for naming this new group as the CL group.

5.3.2. Other meteorites previously proposed to belong to the Coolidge-Loongana grouplet

There are some other ungrouped carbonaceous chondrites, which may belong to the Coolidge-Loongana grouplet. However, as will be argued below, none of these samples share the characteristic properties of the five CL chondrites of this study. Nevertheless, future determination of Ti and Cr isotopic compositions of these meteorites will probably help to clarify their relationship to the CL chondrites and other groups of carbonaceous chondrites.

5.3.2.1. NWA 779. Weckwerth et al. (2001) proposed that NWA 779 belongs to the Coolidge-Loongana grouplet.
However, the determined mean Fa concentration in olivine is 21 mol%, very different from the Fa values of Loongana (12.5 mol%), Coolidge (14.5 mol%) and the other samples described by us (12.8–14.7 mol%; Table 4). Moreover, several volatile element concentrations are considerably higher in NWA 779. These are, for example K₂O (0.08 wt.% in NWA 779 compared to 0.01–0.02 wt.% in the CLs; Table 9), MnO (0.20 wt.% compared to 0.14–0.16 wt.%; Table 9), and Zn (110 ppm compared to 33–57 ppm; Table 10). A further obvious difference is the low concentration of Ni and Co in NWA 779 (6000 and 300 ppm, respectively), compared to 12,800–13,900 ppm and 560–600 ppm, respectively, in the five investigated CL meteorites (Table 9). This is difficult to explain by terrestrial weathering, since most of the CL samples are also strongly weathered. Obviously, NWA 779 contains (as a primary feature) much lower amounts of Fe-Ni metal than the CL meteorites. For these reasons, NWA 779 does not belong to the proposed CL group, but appears to be correctly classified as CV3 (of high petrologic type) in the MBDB.

5.3.2.2. Hammada al Hamra (HaH) 073. Weckwerth and Weber (1998) and Weckwerth (2014) conclude that Hammada al Hamra (HaH) 073, currently classified as C4 (MBDB), belongs to the Coolidge-Loongana grouplet. The O isotopic composition of this sample plots within the range of CV-CK chondrites on the CCAM line in δ¹⁸O-δ¹⁷O space (MBDB). Its mean Fa content in olivine is 17.9 mol%, i.e., outside the range of the five investigated CL samples (12.5–14.7 mol%) and it contains considerably more (~33 vol%) fine-grained matrix (Weckwerth and Weber, 1998) than the here studies samples (17–21 vol%; Table 2). The mean apparent chondrule size of HaH 073 is 580 μm (Weckwerth and Weber, 1998), also outside the range of the measured CL values (404–477 μm; Table 7). The concentrations of Ni and Co in HaH 073 (12,600 and 575 ppm, respectively; Choe et al., 2010) are slightly lower than those in the CL samples (12,800–13,900 ppm and 560–600 ppm, respectively; Table 9), indicating a slightly lower abundance of Fe-Ni metal. The concentrations of many volatile elements in HaH 073 are considerably higher than in CL chondrites. The Na₂O concentrations in the latter are 0.18–0.32 wt.% (Table 9), compared to 0.51 (Weckwerth and Weber, 1998) and 0.50 (Choe et al., 2010), respectively, in HaH 073. The same is true for K, which shows low concentrations of 0.01–0.02 wt.% in CL chondrites (Table 9) and high concentrations between 0.034 (Choe et al., 2010) and 0.040 wt.% (Weckwerth and Weber, 1998) in HaH 073. The MnO concentration in HaH 073 (0.17 wt.%; Weckwerth and Weber, 1998; Choe et al., 2010) is somewhat higher than in the CL samples (0.14–0.16 wt.%), and the Zn concentration is distinctly higher, namely 63–66 ppm (Weckwerth and Weber, 1998; Choe et al., 2010), compared to 33–57 ppm in our samples. However, Weckwerth (2014) revised several HaH 073 values from an earlier publication (Weckwerth and Weber, 1998) and some of the new results are closer to the range of CL values, e.g., MnO (0.13 wt.%) and Zn (20 ppm). However, the new data for Na₂O (0.43 wt.%) and K₂O (0.037 wt.%) are still much higher than those in the CL chondrites. Taking the above differences together, it appears that HaH 073 may be related to, but is not a member of the CL chondrite group.

5.3.2.3. Sahara 00182. Weckwerth (2014) argued that Sahara 00182 (C3-ung; MBDB) belongs to the Coolidge-Loongana grouplet. This meteorite with a mean Fa value of 7.8 mol% (Smith et al., 2004) is probably paired with Sahara 00177 (C3/4-ung; MBDB), which shows a mean Fa value of 7.6 mol%. These Fa values are far outside the range for CL chondrites (12.5–14.7 mol%). Due to the absence of phyllosilicates, Weisberg (2001) classified Sahara 00182 as the first CR3 chondrite. However, its oxygen isotopic composition plots on the CCAM line in δ¹⁸O-δ¹⁷O space (MBDB), which excludes a relationship to CR chondrites. A mean apparent chondrule size of 1010 μm was determined by Smith et al. (2004), a much higher value compared to the CL samples (404–474 μm; Table 7). The modal abundance of matrix is ~28 vol% (Smith et al., 2004), a value which is outside the observed range in CL chondrites (17–21 vol%; Table 2). The concentration of Ni and Co in Sahara 00182 (14,900 and 691 ppm, respectively; Choe et al., 2010) are higher than CL values (12,800–13,900 ppm and 560–600 ppm, respectively; Table 9), indicating a higher abundance of Fe-Ni metal. Furthermore, Na₂O and K₂O concentrations in Sahara 00182 are higher than in the CL chondrites, namely 0.29–0.36 and 0.03 wt.%, respectively (Choe et al., 2010; Weckwerth, 2014), compared to 0.18–0.27 wt.% and 0.01–0.02 wt.% (Table 9). Due to these differences it is improbable that Sahara 00182 is closely related to the CL chondrites. The strongest argument against such a relationship is provided by the distinctly different chemical composition of matrix. The SiO₂ and MgO concentrations in the matrix of Sahara 00182 are 24.2 and 21.7 wt.%, respectively, compared to 30.3 and 28.3 wt.% in the CL chondrites (Table 8). These values from Smith et al. (2004) have been recalculated to totals of 95 wt. % for comparison. There are even more extreme differences in the matrix concentrations of CaO and S. While the matrix in CL chondrites is characterized by very low concentrations of CaO and S (0.97–1.61 wt.% and 0.04–0.35 wt.% respectively; Table 8), matrix in Sahara 00182 contains 5.75 wt. % CaO and 2.50 wt.% S (Smith et al., 2004).

5.3.3. Anticipated characteristics of unmetamorphosed, yet to be detected samples

Since all investigated CL meteorites are affected by thermal metamorphism (petrologic types 3.9 to 4; Table 2) the question remains how unmetamorphosed samples might look like. It cannot be ruled out that such samples are hidden among the tangled mass of CV₂₃₄ and CR chondrites from Northwest Africa (NWA) and other dense collection areas. On a first glance, those samples would appear unusual due to their very high chondrule abundances and the correspondingly low amounts of fine-grained matrix (Table 2). Depending on the weathering grade, they would more or less stand out against CV₂₃₄ chondrites by their high metal content and high magnetic susceptibility (Table 3). The O isotopic compositions would plot on the CCAM line, within the range of CV-CK chondrites or to
the direction of lighter isotopic compositions (Fig. 12). In the end, unmetamorphosed CL chondrites would be definitely uncovered by their characteristic depletions of volatile lithophile and chalcophile elements (Figs. 9 and 10), and their Ti and Cr isotopic compositions (Fig. 12b).

5.4. Relationship between CL chondrites and CV and CK chondrites

The recognition of a further distinct carbonaceous chondrite group raises questions about how the characteristics of each of these groups was initially established. Oxygen and Cr isotopic data suggest a relationship between the CL chondrites and CV and CK chondrites. Perhaps their parent bodies formed in a similar nebular region (heliocentric distance) distinct from those of the other carbonaceous chondrite groups. Compared to CV and CK asteroids, the CL asteroid(s) obviously formed from a distinct mix of components, with higher amounts of chondrules and lower amounts of CAIs and matrix. The low matrix content may indicate a correspondingly low amount of co-accreted water ice which led to more reducing conditions during thermal metamorphism.

However, most CV chondrites are of low petrologic type (3.1–3.6; e.g. Bonal et al., 2006), which is in strong contrast to CK and CL chondrites. Most CK chondrites are of petrologic types 4 to 6 and all yet identified members of the CL group are of petrologic types 3.9 to 4 (Table 2). One explanation for these differences is a possible bias of material delivery to Earth by secondary parent bodies, which are in a suitable resonance position just by chance. Greenwood et al. (2020) discuss the relationship between primary and secondary parent bodies, where the latter derived from the former through impact break-up. By this process, secondary bodies form, which do not fully represent the material of the primary body. Even if all primary bodies would have had a similar, for example, onion shell structure of petrologic types (type 6 in the interior, types 5–3 in the outer layers), secondary fragments would mostly consist of material with a restricted range of petrologic types. These types would be overrepresented in our collections, if such secondary parent bodies are the main source of a given meteorite group (e.g., CL chondrites) due to their favorite resonance position.

6. SUMMARY AND CONCLUSIONS

The five ungrouped carbonaceous chondrites Coolidge, Loongana 001, LoV 051, NWA 033, and NWA 13400 can be grouped together to establish the new Loongana (CL) group of carbonaceous chondrites. The results of this study show that CL chondrites are characterized by:

1. Lithophile element ratios (e.g., Al/Mg, Si/Mg) within the range of other carbonaceous chondrite groups
2. Fe-Ni metal abundances considerably higher than for CV, but similar to CR chondrites
3. Chondrule size-frequency distributions similar to CV, but dissimilar to CR chondrites
4. Mean CAI abundances of ~1.4 vol%, i.e., lower than in CV, but much higher than in CR chondrites
5. Very low amounts of matrix (17–21 vol%), the lowest among the main carbonaceous chondrite groups (CI, CM, CO, CV, CR, CK)
6. Nearly equilibrated olivine with mean fayalite (Fa) values between 12.5 mol% (Loongana) and 14.7 mol% (NWA 13400) as a metamorphic effect
7. Lower Al₂O₃ and higher MgO and Cr₂O₃ concentrations in matrix compared to matrix in CV, CK, and CR chondrites, as a possible metamorphic effect
8. Considerable bulk depletion of volatile elements (Mn, Na, K, Rb, Cs, Zn, Se, Te, Pb, Tl) compared to all other main carbonaceous chondrite groups, reflecting the very low matrix abundance.
9. Bulk O isotope compositions plotting along the CCAM line (Δ¹⁷O = −3.96 to −5.47‰), partly overlapping with the CV and CK chondrite fields but including samples that are ¹⁶O-enriched
10. Unique positions in the e⁴⁸Cr–e⁵⁰Ti isotope plot, with e⁴⁸Cr values similar to CV, CK, and CO, but e⁵⁰Ti values similar to CR chondrites.

Loongana-type (CL) chondrites show many similarities to CV and CK chondrites and may have accreted from a similar nebular reservoir, which, however, was characterized by much lower matrix and CAI abundances. The CL chondrites also display some similarities to CR chondrites (e.g., average magnetic susceptibility), but these groups are not related, as is evident from their different O and Cr isotopic compositions (Figs. 11 and 12a), and their different frequency distributions of apparent chondrule sizes (Table 7, Fig. 5).

One of the remaining key questions is whether unidentified or misidentified CL members of lower petrologic types are hidden in our collections, possibly among the large number of CVred and CR chondrites from hot and cold deserts. Such samples would be important to investigate the pristine state of these meteorites and their components.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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APPENDIX A. SUPPLEMENTARY MATERIAL

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