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A Mid-Infrared Study of Synthetic Glass and Crystal Mixtures Analog to the Geochemical Terranes on Mercury --Manuscript Draft--

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Abstract:	The MERTIS (MErcury Radiometer and Thermal Infrared Spectrometer) onboard of the BepiColombo ESA/JAXA mission to Mercury will map the surface of Mercury in the wavelength range of 7-14 µm and for the interpretation of these spectra a database of analog materials is needed. We analysed bulk grain size fractions of a series of analog materials relevant to the distinct terranes of Mercury in diffuse reflectance in the mid-infrared (2.5 µm to 18 µm). Mineral mixtures cover a wide range of modal amounts of forsterite, enstatite, diopside and plagioclase, the resulting spectra can be divided into three distinct groups: (1) is dominated by a single glass feature, (2) by forsterite bands, and (3) by pyroxene bands. Despite often high contents, plagioclase features, are usually 'overprinted' by forsterite and pyroxene bands. Spectral parameter CF, an easy obtainable proxy for chemistry (SiO2) and polymerization (SCFM) places the hermean mixtures mostly in the intermediate and basaltic range. The correlation of parameters easily obtainable in remote sensing, Mg/Si ratio, and CF, allows differing materials from high-energy evaporation processes in impacts from such formed in igneous processes. Preliminary comparison with a spectrum covering most of the hermean surface shows some similarity with band positions of the Inter Crater Plain and Heavily Cratered Terrains (IcP-HCT) and High-Mg Northern Volcanic Plains (High-Mg NVP) mixtures, but none of our spectra is able to reproduce the remote sensing data entirely.						
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Abstract

33	The MERTIS (MErcury Radiometer and Thermal Infrared Spectrometer) onboard of the BepiColombo
34	ESA/JAXA mission to Mercury will map the surface of Mercury in the wavelength range of 7-14 μm and
35	the interpretation of these spectra a database of analog materials is needed. We analysed bulk grain size
36	fractions of a series of analog materials relevant to the distinct terranes of Mercury in diffuse reflectance
37	in the mid-infrared (2.5 μm to 18 μm). Mineral mixtures cover a wide range of modal amounts of
38	forsterite, enstatite, diopside and plagioclase, the resulting spectra can be divided into three distinct
39	groups: (1) is dominated by a single glass feature, (2) by forsterite bands, and (3) by pyroxene bands.
40	Despite often high contents, plagioclase features, are usually 'overprinted' by forsterite and pyroxene
41	bands.
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43	the hermean mixtures mostly in the intermediate and basaltic range. The correlation of parameters
44	easily obtainable in remote sensing, Mg/Si ratio, and CF, allows differing materials from high-energy
45	evaporation processes in impacts from such formed in igneous processes.
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48	Northern Volcanic Plains (High-Mg NVP) mixtures, but none of our spectra is able to reproduce the
49	remote sensing data entirely.
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55 1. Introduction

57 BepiColombo will map the surface of Mercury in the wavelength range of 7-14 μ m, with a spatial 58 resolution of around 500 meters (Hiesinger et al., 2020). Spectral infrared features permit the direct 59 determination of Mercury's surface mineralogy (e.g., Rothery et al., 2020), which is not possible with 60 ground-based observations at this resolution (e.g., Sprague et al., 2009). In order to extract a more 61 accurate, as well as spatially-resolved mineralogy from the future spectral data, laboratory studies for 62 comparison with BepiColombo measurements are required to support spectral modelling. In order to 63 achieve this, the IRIS laboratory in Münster and the BED laboratory in Berlin were installed to 64 characterize a wide range of samples relevant to Mercury (e.g. Weber et al. 2018). 65 There are no known meteorites from Mercury (e.g., Weber et al., 2016). Hence, we rely on 66 remote sensing observations to study the mineralogy of Mercury. Multispectral imaging with the 67 Mercury Dual Imaging System (MDIS) and spectral reflectance measurements with the Mercury 68 Atmospheric and Surface Composition Spectrometer (MASCS) onboard MESSENGER recognized the 69 occurrence of graphite (Peplowski et al., 2016), sulphides (Vilas et al., 2016), and ice in polar regions 70 (Neumann et al. 2013). Spectral measurements also show that the crust of Mercury contains Fe²⁺-poor 71 silicate minerals (Izenberg et al., 2014). However, spectral data from MESSENGER were insufficient to 72 identify the mineralogy of silicate phases and their abundances. Additionally, information on the 73 chemical composition of the hermean surface is based on data from X-ray, gamma-ray, and neutron 74 spectrometers onboard the MESSENGER spacecraft (e.g., Peplowski et al., 2011, 2016; Nittler et al., 75 2020). Major element ratios obtained from the various instruments allowed the identification of several 76 chemically different regions (the so-called 'terranes') on the hermean surface. The terranes were named 77 the Low-Mg Northern Volcanic Plains (Low-Mg NVP), the High-Mg Northern Volcanic Plains (High-Mg

The mid-infrared spectrometer MERTIS (MErcury Radiometer and Thermal Infrared Spectrometer) of

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78 NVP), the Smooth Plains, the Inter Crater Plains and Heavily Cratered Terrains (IcP-HCT), and the High-79 Mg Province (Nittler et al., 2011; Peplowski et al., 2016; Weider et al., 2015; vander Kaaden et al., 2016; 80 Peplowski and Stockstill-Cahill, 2019). Typical rock types on Mercury's surface can broadly be described 81 as Fe-free and S-enriched komatiites, boninites, and andesites (e.g., Weider et al., 2015; Vander Kaaden 82 et al., 2016; Peplowski and Stockstill-Cahill, 2019). Thermodynamic modeling and experimental petrology 83 provide first order constraints on the mineralogy of such rocks, which are likely to be dominated by 84 olivine, pyroxenes, plagioclase ± quartz, and (Mg,Ca)S sulfide (e.g., Charlier et al., 2013, Stockstill-Cahill 85 et al., 2012; Namur & Charlier 2017; Vander Kaaden et al., 2017; Renggli et al., 2022). Data from midinfrared spectroscopy from terrestrial telescopic observations integrating large surface regions also 86 87 indicate a hermean mineralogy dominated by plagioclase and pyroxene (Donaldson-Hanna et al., 2007; 88 Sprague et al., 1994, 2000, 2002, 2007; Sprague and Roush, 1998; Emery et al., 1998; Cooper et al., 89 2001).

90 In this study, we performed spectroscopic measurements in the mid-infrared wavelength range 91 of analog glass and crystal mixtures relevant to Mercury's surface. The compositions of the glasses and 92 crystals were chosen based on results from high-temperature crystallization experiments (1480-1100°C), 93 that were equilibrated at crustal pressure (1 kbar) and reduced redox conditions (around IW-5; Namur 94 and Charlier, 2017). The bulk compositions of these experiments are representative of the five main 95 chemically distinct regions of the hermean surface introduced above, and the quenched products 96 contain some of the following phases: silicate glass, plagioclase, forsterite, diopside, enstatite, quartz, 97 FeSi metal, and sulfide (i.e., stoichiometric FeS). The samples we measured in this study are mixtures of 98 pure endmember minerals and glass, very similar to the chemical compositions of the experimental 99 products presented by Namur and Charlier, 2017. Synthetic glasses used in this study were previously 100 described in detail in Morlok et al. (2021).

101 The overall aim of this study is to (a) provide mid-infrared reflectance spectra of hermean 102 mineral/glass mixtures in different size fractions to allow granumeletric studies, and under varying 103 observational geometries to accommodate for changing observational conditions in orbit, and (b) 104 provide first spectral parameters for the comparison with future data obtained by MERTIS.

	ID	mineral / features	locality (origin)								
	13	Quartz	Mongolia								
	22	Diopside	Otter Lake, Quebec, Canada								
	28	Labradorite	Ihosy, Madagascar								
	53	Enstatite	Odegardens Verk, Bamble, Norway								
	249	Olivine	Dreiser Weiher, Germany								
106	Table	1: Sources of natural crystalline phases us	sed in this study. Samples are from larger,								
107	crushe	ed single crystals and handpicked afterwa	rds. Spectra can be found in Figure 2f. Chemical								
108	data s	ee Table 3.									
109											
110	2. Sam	ples and Techniques									
111	2.1. Sa	mples									
112	Compo	ositional and mineralogical characteristics of f	ive terranes are covered in this study (Peplowksi et								
113	al., 20	., 2015, Weider et al., 2015; Namur and Charlier, 2017; Denevi et al., Nittler et al., 2020). Large lava									
114	flows (~3.5 Ga) are typical of the Northern Volcanic	Plains (Low-Mg and High-Mg NVP). The Low-Mg NVP								
115	is char	acterized by low Mg/Si ratios and a wider ran	ge of Al contents. This results in high-plagioclase								
116	abund	ance, together with diopside and forsterite, w	vhereas the High-Mg NVP lavas contain a in								
117	compa	rison higher olivine abundance.									

The intermediate Inter Crater Plain and Heavily Cratered Terrains (IcP-HCT), characterized by medium
 sized craters, show high Al contents, and contrasting low Mg. This resulted in high modal amounts of
 plagioclase.

3.5 to 3.9 Ga old Smooth plains cover ~40% of the hermean surface and are characterized by flat plains
with numerous wrinkle ridges. The chemistry shows lower Mg/Si, S/Si, and Ca/Si ratios compared to the
other regions, resulting in a mineralogy dominated by plagioclase, but minor mafic components. The old
(4.2 - 4 Ga) High-Mg Province is characterized by high Mg/Si and Ca/Si, and low Al/Si ratios. The samples
used in this study reflect a relatively low a a series of natural crystalline materials (Table 1). We selected
a series of modal (wt%) phase compositions for each surface region identified in Namur and Charlier
(2017) (Fig. 1). Chemical bulk compositions of the mixtures were calculated using the chemical

128 composition of the starting materials and the individual weighed phase abundances (Table 2).

129 Detailed information about the minerals labradorite, enstatite, and forsterite (IDs 28, 53, 249) used in

this study are given in Weber et al. (2021) and Reitze et al. (2021). ID 22 is a diopside, crushed to grain

sizes from 25 μ m – 250 μ m which shows no sign of impurity. However, in the grain size fraction from 0

 $\mu m - 25 \mu m$ we observe signs (<5%) of serpentinization (van der Meer et al., 1995; Daly et al., 2011).

However, this has only small impact on the overall bulk compositions of our samples because we usedlow proportions of ID 22 Diopside in the mixtures.

The mineral samples are natural terrestrial materials. While the chemical composition of these minerals is, overall, similar to minerals on Mercury, these phases formed in vastly different environments. In particular, terrestrial magmatism is much more oxidized than expected for Mercury, with the latter of which is characterized by redox conditions buffered by the iron-wüstite buffer (IW-7 to IW-3), whereas magmatism on the Earth usually happens at redox conditions between IW-2 to IW+8 (McCubbin et al., 2012; Namur et al., 2016a,b; Cartier and Wood, 2019). As a result, for example different Fe^{2+/}Fe³⁺ ratios in olivine and pyroxene can be expected. 142 Silicate glasses were synthesized to represent the amorphous component in our mixtures. Details on the

synthesis protocol can be found in Morlok et al. (2021). We produced a series of six synthetic glasses

using reagent grade oxides as starting materials. Melts were kept in a gas mixing furnace for 6 h at

145 1450°C at the graphite-CO buffer (IW-6.7). The glasses cover the observed compositional range of the

- 146 terranes (e.g., 1.6 wt% 19.0 wt% MgO).
- 147 Size fractions for each mixture were mixed separately (<25 μm, 25 μm 63 μm, 63 μm -125 μm, 125 μm
- $-250 \ \mu m$). This results in minor divergences from the modal starting compositions based on the
- 149 laboratory experiments (see Table 2). The finest fraction (<25 μm) was chosen to represent the regolith
- 150 grain size, which exhibits the important Transparency Feature (TF), a characteristic reflectance low. The
- 151 other size fractions were selected to cover intensity changes related with grain sizes. Increasing grain size
- is correlated with increasing intensity (e.g., Salisbury, 1993; Mustard and Hayes, 1997).
- 153 For measurements, the samples were filled into aluminum sample cups, and the surface flattened with a
- 154 spatula (see Mustard and Hayes, 1997 for details).
- 155
- 156 Figure 1: Overview of the modal composition of the selected samples (Namur and Olivier, 2017). (a)
- 157 pyroxene, plagioclase, and forsterite, (b) pyroxene, forsterite, and glass. All in wt%.



159 2.2. Infrared Studies

160 We used a Bruker Vertex 70v infrared spectrometer with a liquid nitrogen cooled MCT (HgCdTe) detector 161 at the IRIS laboratory in Münster. Analyses were made in the wavelength region of 2.5 μ m to 18 μ m in 162 low vacuum (100 Pa). For each spectrum, 512 scans were accumulated at a spectral resolution of 4 cm⁻¹ 163 (20 nm at 7 μ m, 80 nm at 14 μ m). This spectral resolution is comparable to MERTIS onboard 164 BepiColombo (e.g., 90 nm, Hiesinger et al. 2010 and 2020). For calibration, the instrument background was eliminated by ratioing the sample spectra against the spectrum of a diffuse gold standard 165 166 (INFRAGOLDTM) measured under the same conditions. Analyses were made at 13° incidence (i) and 13° 167 emergence (e), 20° (i)/30°(e), and 30°(i)/30°(e) in order to simulate observational geometries of an 168 orbiter. The results presented in the main text were made at 20°(i) and 30°(e), results for the other 169 settings are available in the supplement. The results are presented in reflectance from $6 \,\mu\text{m} - 18 \,\mu\text{m}$ 170 (Fig. 2a-e), which covers the range of the MERTIS instrument (7 μ m – 14 μ m) but omits volatile features. 171 It also includes the Christansen Feature (CF, the Reststrahlenbands RB; the molecular vibration modes of 172 the materials), and the Transparency Feature (TF; typical for the finest <25 μ m size fractions). 173 Since the aim of our study is to compare the laboratory data with future remote sensing data, 174 measurements from the laboratory have to be recalculated from reflectance to emissivity using 175 Kirchhoff's law: $\varepsilon = 1 - R$ (R = Reflectance, $\varepsilon =$ Emission, Nicodemus, 1965). However, a prerequisite for the validity of Kirchhoff's law is that the reflected light should be collected in all directions. In an ideal 176 177 case directional emissivity and directional hemispherical reflectance should be compared. However, we 178 used a bi-directional setup. Earlier studies showed that this approach mainly affects the reflectance 179 intensity, but not band positions. This must however be kept in mind when comparing the results in a 180 quantitative manner with emission data (Salisbury et al., 1991; Hapke, 1993; Thomson and Salisbury, 181 1993; Salisbury et al., 1994; Christensen et al., 2001). The spectra presented in this study are accessible 182 on our online database at the Institut für Planetologie in Münster (http://bc-mertis-pi.uni-muenster.de/).

	Low- Mg NVP		High- Mg NVP			Smoot h Plains			IcP- HCT			
	ID349 (Y131)	ID 350 (Y172) P	ID 355 (Y133)	ID 356 (Y143)	ID 357 (Y144)	ID 345 (Y140)	ID 344 (Y143)	ID 343 (Y144)	ID 351 (Y126)	ID 352 (Y131)	ID 353 (Y132)	ID 354 (Y146)
Glass	86.9	29.5	96.8	42.0	30.2	90.7	53.8	42.3	78.6	66.6	42.3	20.6
Forster	8:2	14.3	3.2	12	10.7	9.3	5.1	5.9	21.4	29.7	32.5	31.5
ite Diopsi de	4.9	17.1		9.1	12.1		2.2	10.7		3.7	8.1	8.3
Enstati te				16.0	20.3		11.3	7.4				5.0
Plagioc lase		39.1		20.8	26.6		27.6	33.7			17.1	33.4
Quartz												1.2
SiO2	64 52	59 35	58 71	59 01	60 41	62 10	61 34	59 76	54 85	59.00	55 73	55 72
TiO2	0.20	0.09	0.40	0.11	0.10	0.42	0.13	0.11	0,28	0.16	0.11	0.07
AI2O3	13.42	15.40	10.61	12.34	12.04	12.97	16.02	15.96	12,98	10.29	11.31	12.47
Cr2O3	0.00	0.01	0.00	0.01	0.01	0.00	0.01	0.01	0,01	0.01	0.02	0.02
FeO	0.97	1.79	0.32	2.80	3.20	0.82	1.63	1.56	1,89	2.70	3.03	3.42
MnO	0.10	0.06	0.17	0.07	0.05	0.16	0.07	0.07	0,08	0.10	0.09	0.06
MgO	9.03	10.57	19.99	14.86	14.68	12.83	9.20	9.22	20,56	18.74	19.78	19.28
CaO	6.25	8.89	6.94	6.91	6.41	6.18	6.53	8.63	5,64	4.79	6.27	5.95
Nazo	5.22	3.92	3.19	3.64	3.29	4.36	4.70	4.34	3,84	4.01	3.46	3.06
Total	99.91	100 31	100.43	99 93	100 39	100.02	99.85	99.86	100.24	99 97	99 94	100 24
Total	55.51	100.51	100.45	55.55	100.55	100.02	55.85	55.00	100,24	55.57	55.54	100.24
	High- Mg Prov.			Crystal line					Glass			
	ID 348 (Y126)	ID 347 (Y131)	ID 346 (Y146)	ID 249 Forste rite	ID 22 Diopsi de	ID 53 Enstati te	ID 28 Plagio clase	ID 13 Quarz	ID339	ID 338	ID 181	ID 174
Glass	70.5	42.5	27.7						100	100	100	100
Forster ite	29.5	36.2	35.8	100								
Diopsi de		21.2	25.3		100							
Enstati te			1.3			100						
Plagioc lase			7.7				100					
Quartz			2.0					100				
SiO2	53.87	55.14	56.74	40.96	55.08	57.48	55.63	101.42	58.62	55.90	59.29	64.26
TiO2	0.30	0.11	0.09	0.02	0.05	0.04	0.01	0.01	0.35	0.41	0.41	0.46
Al2O3	7.73	6.63	6.41	0.04	0.31	0.14	27.87	0.01	16.49	7.31	10.95	14.30
Cr2O3	0.01	0.02	0.02	0.04	0.01	0.01	0.02	0.01	0.00	0.00	0.00	0.00
FeO	2.51	3.67	3.90	8.38	2.63	9.09	0.10	0.01	0.13	0.10	0.05	0.05
MnO MaQ	0.16	0.11	0.09	0.11	0.14	0.04	0.01	0.01	0.08	0.25	0.17	0.16
	28.30	23.8/ 782	23.25 7 71	50.50 0.00	10.8U 25 10	33.10 0.21	10.01	0.00	12.43 7 15	20.94 7 15	19.00 7.16	8.97 6.81
Na2O	2.33	2.59	2.16	0.09	0.19	0.02	5.29	0.00	4.88	2.95	3.29	4.81
K20	0.08	0.10	0.13	0.00	0.01	0.01	0.31	0.01	0.14	0.11	0.11	0.19
Total	100.35	100.06	100.51	100.14	100.30	100.21	99.57	101.50	100.27	101.11	100.44	100.01

185	Table 2. Chemical bulk compositions (wt. %) and modal mixtures used in this study. Bulk chemical
186	compositions of the mixtures used in this study (in wt.%) were calculated using the chemical
187	compositions of the components and the modal compositions. ID + Number: Sample ID used in the IRIS
188	database (see 2.2. Infrared Studies). Name of sample based on starting composition used following
189	Namur and Charlier (2017). Fo#: Forsterite content, En#: Enstatite content, An#: Anorthite content, Wo#:
190	Wollastonite content (mol.%).
191	
192	3. Results
193	Most samples show spectral features at wavelengths shorter than 7 μ m, usually at 2.7 μ m – 3.0 μ m,
194	3.4 μ m– 4 μ m, and 5 μ m – 7 μ m. These are volatile features due to adsorption and absorption of
195	terrestrial water and unavoidable minor contaminations of the terrestrial minerals used for the mixtures.
196	
197	3.1. Low-Mg Northern Volcanic Plains
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197 198 199	3.1. Low-Mg Northern Volcanic Plains The glassy end member is represented by low Mg sample ID 174 (Morlok et al., 2021), and has the CF at 7.9 μ m, the TF at 11.8 μ m, and a single RB at 9.6 μ m. The spectrum of ID 350 has the CF between 7.8 μ m
197 198 199 200	3.1. Low-Mg Northern Volcanic Plains The glassy end member is represented by low Mg sample ID 174 (Morlok et al., 2021), and has the CF at 7.9 μm, the TF at 11.8 μm, and a single RB at 9.6 μm. The spectrum of ID 350 has the CF between 7.8 μm and 8.1 μm, with several pyroxene RB at 9.4 μm, 9.9 μm, 10.2 μm, 10.5 μm – 10.6 μm and 10.8 μm. The
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197 198 199 200 201 202 203 203 204	3.1. Low-Mg Northern Volcanic Plains The glassy end member is represented by low Mg sample ID 174 (Morlok et al., 2021), and has the CF at7.9 μm, the TF at 11.8 μm, and a single RB at 9.6 μm. The spectrum of ID 350 has the CF between 7.8 μmand 8.1 μm, with several pyroxene RB at 9.4 μm, 9.9 μm, 10.2 μm, 10.5 μm – 10.6 μm and 10.8 μm. TheTF is located at 11.3 μm – 11.5 μm (Tab.4; Fig.2a, f). The dominance of the glassy component results in asimpler spectrum with few bands for ID 349: the CF is at 7.8 μm - 7.9 μm, the TF at 11.6 μm. Thestrongest olivine RB is at 9.4 μm -9.5 μm, with minor features at 10.2 μm and 10.5 μm - 10.6 μm (Tab.4;Fig.2a, f).
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197 198 199 200 201 202 203 204 205 206	3.1. Low-Mg Northern Volcanic Plains The glassy end member is represented by low Mg sample ID 174 (Morlok et al., 2021), and has the CF at 7.9 µm, the TF at 11.8 µm, and a single RB at 9.6 µm. The spectrum of ID 350 has the CF between 7.8 µm and 8.1 µm, with several pyroxene RB at 9.4 µm, 9.9 µm, 10.2 µm, 10.5 µm – 10.6 µm and 10.8 µm. The F is located at 11.3 µm – 11.5 µm (Tab.4; Fig.2a, f). The dominance of the glassy component results in a simpler spectrum with few bands for ID 349: the CF is at 7.8 µm - 7.9 µm, the TF at 11.6 µm. The strongest olivine RB is at 9.4 µm -9.5 µm, with minor features at 10.2 µm and 10.5 µm - 10.6 µm (Tab.4; Fig.2a, f).
197 198 199 200 201 202 203 204 205 206 207	3.1. Low-Mg Northern Volcanic Plains The glassy end member is represented by low Mg sample ID 174 (Morlok et al., 2021), and has the CF at 7.9 µm, the TF at 11.8 µm, and a single RB at 9.6 µm. The spectrum of ID 350 has the CF between 7.8 µm and 8.1 µm, with several pyroxene RB at 9.4 µm, 9.9 µm, 10.2 µm, 10.5 µm – 10.6 µm and 10.8 µm. The TF is located at 11.3 µm – 11.5 µm (Tab.4; Fig.2a, f). The dominance of the glassy component results in a simpler spectrum with few bands for ID 349: the CF is at 7.8 µm - 7.9 µm, the TF at 11.6 µm. The strongest olivine RB is at 9.4 µm -9.5 µm, with minor features at 10.2 µm and 10.5 µm - 10.6 µm (Tab.4; Fig.2a, f).

209 3.2. High-Mg NVP

- The glass-rich samples ID 355 and pure glass ID 181 have the CF at 8 μm to 8.1 μm and the TF at 11.9 μm.
- They have a strong RB at 9.7 μm to 9.8 μm, ID 355 also has weak olivine bands at 10.5 μm. Crystal-
- bearing ID 356 and ID 357 show similar pyroxene band positions, although they are different in
- 213 intensities and band shapes. Their CF is between 7.8 μm and 8.2 μm (Table 4; Fig. 2b, f). Common
- 214 pyroxene RBs are at 9.4 μm -9.5 μm, 10.2 μm, 10.5 μm, 10.8 μm, and 11.3 μm. At longer wavelengths,
- 215 bands occur at 14.4 μm-14.5 μm, and 15.6 -15.8 μm. The TF is located at 11.4 μm.
- 216

217 **3.3 Intra Crater Plains - Heavily Cratered Terrane (IcP-HCT)**

The glass endmember, represented by high-Mg glass ID 181 (Morlok et al., 2021) has the CF at 8 μm to
8.1 μm, the TF at 11.9 μm, and a single RB at 9.8 μm. Similar to the High-Mg Province, the spectra of the
IcP-HCT region (ID 351, ID 352, ID 353, ID 354) are dominated by olivine features, reflecting the forsterite
modes ranging from 21 to 33 wt% (Table 4; Fig. 2c, f). The CF is located between 7.8 μm and 8.2 μm. The
strongest RBs are at 9.4 μm - 9.7 μm, 10.2 μm, 10.5 μm-10.6 μm, and 11.9 μm. The TF is again a broad
feature without distinct maximum. A dip at 7 μm is observed for diopside-rich samples ID 353 and ID
354.

225

3.4. Smooth Plains

- 227 Glass-dominated mixture ID 345 has a CF range from 7.9 to 8 μm, with an RB at 9.6 μm, 10.2 μm and
- 228 10.5-10.6 μm. The TF is located at 11.8 μm.
- 229 The-Smooth Plains samples ID 343 and ID 344 show generally similar pyroxene band positions (Table 4;
- Fig. 2d, f) reflecting their similar bulk composition, but with ID 344 having more pronounced, 'sharper'
- bands. The CF is between 7.8 µm and 8 µm for ID 343, and between 7.8 µm and 7.9 µm for ID 344.
- 232 Significant RBs are between 9.4 μm and 9.6 μm and at 9.9 μm, 10.2 μm, 10.5 μm, 10.8 μm an 11.4°μm.

The TF is located at 11.7 μm. Various minor spectral features are found at longer wavelengths, mainly
from 15.5 μm to 15.8 μm. The diopside-rich sample ID 343 shows a dip at 6.9 μm to 7 μm (or a small
feature at 7.3 μm) in the finest grain size fraction.

236

237 3.5. High-Mg Province

The glassy end member ID 338 has the CF at 8.2 μm, TF at 12.1 μm and the dominating RB at 9.9 μm.

As a result of the high forsterite contents (30-36 wt%), spectra of mixtures ID 346, ID347, and ID348 are

quite similar (Table 2; Fig. 2e, f). The plagioclase and pyroxene contents show no significant bands in the

spectra. The CF ranges from 7.8 μm to 8.6 μm, and the strongest RB is at 10.6 μm. Further common

bands are found at 10.2 μm and 11.9 μm – 12 $\mu m.$ ID 346 and ID 347 share further RBs at 9.4 μm and

243 10.8 μm. The TF is a broad feature without a clear peak. Features at longer wavelengths are seen

between 15.8 μm and 16 $\mu m.$ Samples ID 346 and ID 347 shows a dip at 6.9 μm in the finest grain size

245 fractions.

246

Figure 2a-f: Mid-infrared reflectance spectra of the grain size fractions. Black: 125 - 250 μm, dark grey:
63 – 125 μm, grey: 25 - 63 μm, light grey: 0-25 μm. Gl: glass, Fo: forsterite, Di: diopside, Pl: plagioclase,
En: enstatite, Qz: quartz. Vertical green lines: 249 forsterite bands, vertical yellow lines: pyroxene
features (see Figure 3).





Figure 2













255 4. Discussion

256 **4.1. Interpreting the mixtures**

257 Despite the often-complex mixtures of up to five crystalline and one amorphous phase, the resulting 258 spectra of the mixtures can be divided into three groups (Table 3). The simplest spectra are dominated 259 by glass, demonstrated by prominent RB from 7.9 μ m – 8.3 μ m. Besides the 'pure' glassy end members 260 (ID 174, ID181, and ID 338), sample ID 355 High-Mg NVP with only minor (3 wt%) crystalline content 261 shows barely any recognizable additional bands (Fig. 2b). In the second group, forsterite features tend to 262 dominate (Table 3): bands near 9.4 μ m to 9.5 μ m, 10.2 μ m, 10.6 μ m, 11.9 μ m, and 15.9 μ m -16 μ m 263 (Fig.2c, e). In the finest size fraction (< 25 μ m), the TF the 10.6 μ m band is turned into a broader feature 264 with another peak at 11.2 μ m to 11.3 μ m. 265 A third group exhibits the pyroxene RB features of diopside and enstatite near 8.9 µm to 9.1 µm, 9.4 µm 266 - 9.5 μm, 9.9 μm, 10.2 μm, 10.4 μm - 10.8 μm, 11 μm - 11.1 μm - 11.3 μm, and 11.4 μm to 11.6 μm 267 (Fig.2a-e). The finest grain size fractions of diopside-rich samples also show a dip at 6.9 μ m to 7 μ m, 268 which is also prominent in the finest grain size fraction of the ID 22 diopside bulk material. 269 Although plagioclase feldspar is often a dominant component in the mixtures, its RB features tend to be 270 subdued by the forsterite and enstatite bands. The only RB, which is visible in all samples with a high 271 amount of plagioclase is its RB 1 as a shoulder at around 8.3 μ m (e.g. 356 and 357 High-Mg NVP; Fig. 4b). 272 This is probably caused by its flat spectrum, that is dominated by a broad RB at 8.7 µm, and particular at 273 9.9 μ m to 10 μ m and 10.5 μ m to 10.6 μ m, which are superimposed by the more pronounced pyroxene 274 and olivine bands. However, further hints of significant plagioclase contents are higher intensities in the 275 \simeq 10 μ m to 10.5 μ m range due to the strong plagioclase double-feature, which provides a generally broad 276 feature, which serves as some kind of underlying continuum, as observed in samples ID 350 and ID 356 277 (Fig.2a, b). In the wavelength range around 12 μ m glasses and plagioclases have overlapping TF. In 278 earlier experiments using plagioclase (ID28) with varying degrees of Al-Si order (Reitze et al. 2021a) 279 plagioclases with low degree of Al-Si order show a single TF, such with a higher order a double peak or

peak with shoulder, as seen in the case of the plagioclase ID28 used in this study (e.g. 350 Low-Mg NVP
or 357 High-Mg NVP; Fig.2a, b).

282	The quartz content is too low (few wt%) to produce any recognizable RB (e.g., 2 wt.% in ID 346, 1.2 wt.%
283	in ID 354; Fig.4b and e). This raises the question of the 'critical' concentration (i.e. detection limit) of a
284	phase to be recognizable in a mixture. In simple mixtures of glass and a crystalline phase, minute
285	contents of 3 wt.% olivine in ID 355 High-Mg NVP(Fig.2b) provide just enough signal to identify the
286	mineral with features at 10.5 μm and 11.9 μm (Table 4). Contents of 9.3 wt% olivine in ID 345 Smooth
287	Plains (Fig.2d) are easily recognizable with bands at 9.6 μ m, 10.2 μ m, 10.6 μ m, and 11.4 μ m (Table 4).
288	However, two minor components of 8.2 wt.% olivine and 4.2 wt.% diopside in ID 349 Low-Mg NVP
289	(Fig.4a) only allow the clear identification of the olivine 9.4-9.5 μ m, 10.2 μ m, 10.5 μ m-10.6 μ m, and 11.6
290	μ m. Increasing number of phases will make the identification of a minute phase in the crowded range
291	around 10 μm difficult. For the semi-quantitative identification of further components, rationing of
292	spectral parameters could provide an alternative.

	RB Features	Samples
ID 338, 181,174 Glass	9.6 - 9.9 μm	ID 174, ID181, ID 338,
		ID 355 High-Mg NVP
ID 249 Forsterite	9.4 μm- 9.5 μm, 10.2 μm, 10.6	ID 349 Low-Mg NVP,
	μm, 11.9 μm, 15.9 μm -16 μm	ID 346, 347, 348 High-Mg Prov.,
		ID 351, 352 353, 354 IcP-HCT,
		ID 345 Smooth Plains,
		ID 355 High-Mg NVP
ID53 Enstatite/ 29 Diopside	8.9 μm - 9.1 μm, 9.4 μm - 9.5	ID 350 Low-Mg NVP,
	μm, 9.9 μm, 10.2 μm, 10.4 μm-	ID 343, 344 Smooth Plains,
	10.8 μm, 11 μm-11.1 μm - 11.3	ID 356, 357 High-Mg NVP
	μm, 11.4 μm to 11.6 μm	

Table 3. Overview of the groups into which the mixture spectra can be divided based on their dominantfeatures.





- Mercury Mixtures (This study)
- O Mercury Glass Component (Morlok et al., 2021)
- ▲ Bulk △ Micro Synthetic Mercury Glasses (Morlok et al., 2017, 2020a) 🔺 Bulk
- Bulk Synthetic Planetary Glasses (Morlok et al., 2020a)
- □ Micro
- ♦ Impact Rocks and Glasses (Morlok et al., 2016a and b)
- K Glass Basalt Evaporation Experiments (Morlok et al., 2020b)



Figure 3. (a) Correlation of SiO₂ abundance with CF position. Results from this study (red circles) mainly fall into the intermediate area. (b) Comparison of the Mg/Si ratio with the CF. The refractory glasses produced with laser-heating (Morlok et al., 2020b) is easy to distinguish from the samples with hermean and planetary compositions. (c) Comparison of SCFM (SiO₂, CaO, FeO, MgO) index of polymerization to CF position. Blue shaded areas and dashed line mark are from Cooper et al. (2002) for terrestrial rocks.

308 4.2 Correlation of Bulk Composition with Spectral Parameters

309 Christiansen Feature and SiO₂

310 The CF can be extracted even from 'noisy' spectra. Hence any correlation of the CF with other features 311 will help to derive basic information about composition and mineralogy from remote sensing data. A 312 classical method to correlate the average CF of all grain size fractions with chemical composition is the 313 comparison of the SiO₂ content (wt.%) with the CF (Cooper et al., 2002). The comparison of the hermean 314 mixtures in this study with earlier results (Fig.3a) shows that most of the mixtures fall close to the 315 correlation line for terrestrial materials, which is the best fit for results of a range of terrestrial rocks 316 scattering a bit due to compositional variances as indicated by the blue shaded areas (Cooper et al., 317 2002). Furthermore, the results fall into or very close to the field for intermediate rocks.

318

319 Christiansen Feature and Bulk Mg/Si

320 Absolute element abundances are difficult to determine from mid-IR remote sensing data. Hence, we

321 use the element ratio between Mg and Si for a correlation with the CF, as ratios of chemical data of two

- major elements is easier to extract from the space probe data (Fig.3b). In combination with the CF it
- 323 provides a set of parameters relatively easily extracted from remote sensing data.

Here, our results fall along the area of synthetic Mercury composition glasses (Morlok et al., 2017a,

325 2020a). The terrestrial impact glasses and the refractory glass from laser-heated melting experiments

326 (Morlok et al., 2020b) are clearly in a separate field compared to our samples relevant to Mercury. This

separation could help to identify glassy material formed in high-velocity impacts from such produced involcanic processes.

329

330 Christiansen Feature and SCFM

The SCFM index expresses the degree of polymerization based on the SiO₂/SiO₂+CaO+FeO+MgO relation (Salisbury and Walter, 1989). Similar to earlier studies, most of the results plot systematically below the terrestrial correlation line (Fig.4c). This could be related to the varying degrees of polymerization of the mixtures, however the correlation between glass content and SCFM is very weak (R²=0.04) The results show similarities to the intermediate, basic and basaltic terrestrial rocks (Cooper et al., 2002)

336

337 4.3. Comparison with Remote Sensing Data of Mercury

338 Only few mid-infrared spectra are so far available for Mercury. Owing to technical reasons they cover

larger surface regions $(10^4 - 10^6 \text{ km}^2)$ and offer only weak spectral contrast and low signal to noise ratios

340 (Sprague et al., 2007). Therefore, we only use as an example the spectrum with the highest signal to

noise for our comparisons. The spectrum integrates a region at about 210–250° longitude and was

obtained by the Mid Infrared Camera (MIRAC) at the Kitty Peak Observatory (Sprague et al., 2000).

The spectrum recalculated from emission (Fig.4) shows a CF at 8.5 μm, three strong RBs at 9.3°μm,

9.9°μm, and 11°μm. A potential TF is visible at 12.4°μm. We compare it with our results of the 0-25 μm

345 fraction of two mixtures reproducing some of the features.

346 The olivine-rich 347 High-Mg Province mixture reproduces the CF of the hermean spectrum, and also has

an RB at 9.4 µm, near the 9.3 µm feature of the surface spectrum. A broad feature at 11°µm overlaps

348 with a (much narrower) band of the surface spectrum at the same position.

- Plagioclase and enstatite-rich 357 High-Mg NVP mixture has a low-point at 8.5 μm, a strong RB at 9.4°μm
- and a minor RB at 9.9°µm. These features are similar to the CF at 8.5 µm and the second and third RB
- 351 (9.3° μ m and 9.9° μ m) of the hermean sample.



354 Figure 4. Comparison of Plagioclase and enstatite-rich 357 High-Mg NVP, and olivine-rich 347 High-Mg 355 Province mixtures with a Mercury surface spectrum (recalculated from emissivity) (Sprague et al., 2000, 356 2007). Shaded areas mark characteristic features of the hermean spectrum. 357 358 5. Summary & Conclusions 359 We mixed bulk grain size fractions of analog materials relevant to the distinct terranes of Mercury. The 360 powdered samples were analyzed in diffuse reflectance in the mid-infrared (2 μ m – 20 μ m). 361 The resulting spectra can be divided into three groups: (1) such as dominated by a single glass feature, 362 (2) a group with forsterite, and (3) a group dominated by pyroxene bands. Plagioclase features, even 363 when the phase is dominating the composition, are usually 'overprinted' by forsterite and pyroxene 364 bands. 365 The spectral parameter CF in comparison with chemistry (SiO_2) and polymerization (SCFM) places the 366 hermean mixtures mostly in the intermediate and basaltic range, similar to findings based in Peplowski 367 and Stockstill-Cahill (2019). The correlation of parameters Mg/Si ratio and CF, that are easily obtainable 368 in remote sensing, allows to distinguish materials that were formed by high energy evaporation 369 processes in impacts from those formed by melting processes. 370 Preliminary comparison with a high-quality spectrum of the hermean surface (Sprague et al., 2000) show 371 some similarity with band positions of IcP-HCT and High-Mg NVP mixtures, but none of our spectra is 372 able to reproduce the remote sensing data entirely. This is not unexpected since further conditions (e.g. 373 temperature) will also affect spectral properties, and this will have to be addressed by future studies. 374 375 376

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Smooth Plains	CF													
ID 343														
0-25 μm	8.04	9.01	9.63	9.92	10.20	10.54	10.82			11.65			15.69	
25-63 μm	7.93		9.51	9.92	10.23	10.49	10.78		11.27				15.77	
63-125 μm	7.90		9.40	9.91	10.23	10.49	10.78						15.76	
125-250 μm	7.84		9.62	9.89	10.21	10.46							15.69	
ID 344														
0-25 μm	7.91		9.54	9.87	10.18	10.51				11.67		14.43		
25-63 μm	7.89		9.53	9.90	10.18	10.48			11.28		13.75	14.43	15.69	
63-125 μm	7.84		9.51	9.89	10.20	10.48			11.30			14.47	15.65	
125-250 μm	7.83		9.51	9.91	10.22	10.45	10.76	11.06	11.38		13.79	14.47	15.53	
ID 345														
0-25 μm	7.97		9.59		10.16	10.56			11.42	11.75			15.81	16.01
25-63 μm	7.93		9.58		10.17	10.55							15.85	16.08
63-125 μm	7.91		9.61		10.16	10.50							15.78	15.96
125-250 μm	7.89		9.59		10.17	10.56								15.93
ID 174														
0-25 μm	7.89		9.58							11.78				
25-63 μm	7.87		9.58											
63-125 μm	7.91		9.58											
125-250 μm	7.91		9.58											
High-Mg														
Province														
ID 346														
0-25 μm	8.60		9.37		10.19	10.64						14.12	16.03	
25-63 μm	7.91		9.37		10.19	10.57	10.79		11.21	11.92		14.84	15.93	
63-125 μm	7.81		9.36		10.19	10.56	10.80			11.90		14.85	15.83	
125-250 μm	7.77		9.36			10.55	10.81			12.41		14.86	15.80	
ID 347														
0-25 μm	8.53		9.39		10.18	10.64	10.84					14.10	16.08	
25-63 μm	8.03		9.38		10.19	10.57	10.79			11.90			15.91	
63-125 μm	7.87		9.38		10.18	10.57	10.79			11.92			15.81	
125-250 μm	7.83		9.37		10.18	10.56	10.79			11.92			15.81	
ID 348														
0-25 μm	8.21		9.63		10.16	10.64				11.96			15.96	
25-63 μm	8.14		9.56		10.16	10.59				11.92			15.94	
63-125 μm	8.11		9.66		10.18	10.57				11.89			15.98	
125-250 μm	8.08		9.61		10.17	10.56				11.89			15.89	
ID 338														
0-25 μm	8.27			9.90						12.06				
25-63 μm	8.21			9.84										
63-125 μm	8.22			9.90										
125-250 μm	8.20			9.87										

Low-Mg NVP	CF										
ID 349											
0-25 μm	7.88		9.45		10.16	10.56			11.62	15.66	
25-63 µm	7.85		9.40		10.16	10.52				15.79	
63-125 μm	7.80		9.39		10.16	10.48				15.71	
125-250 μm	7.76		9.39		10.15	10.50				15.82	
ID 350											
0-25 μm	8.14	8.36	9.40	9.89		10.62		11.32	11.50	15.73	
25-63 µm	7.86		9.36	9.89		10.52	10.82			15.77	
63-125 μm	7.82		9.36	9.91		10.52	10.81			15.77	
125-250 μm	7.76		9.39	9.89	10.19	10.52				15.93	
ID 174											
0-25 μm	7.89		9.58						11.78		
25-63 μm	7.87		9.58								
63-125 μm	7.91		9.58								
125-250 μm	7.91		9.58								
IcP-HCT	_										
ID 351											
0-25 um	8.15		9.63		10.15	10.61			11.94	16.12	
25-63 um	8.09		9.62		10.15	10.55			11.91	15.95	
63-125 um	8.04		9.68		10.16	10.55				15.95	
125-250 um	8.02		9.57		10.16	10.59			11.92	15.91	
ID 352											
0-25 um	8.14		9.54		10.16	10.64		11.30	11.90	15.92	
25-63 um	7.87		9.48		10.16	10.58				15.91	
63-125 um	7.83		9.44		10.16	10.55			11.91	15.91	
125-250 um	7.79		9.39		10.16	10.55			11.90	15.91	
ID 353			5.65		10110	10100			11.50	10101	
0-25 um	8.15		9.66		10.16	10.62		11.24	11.92	16.01	
25-63 um	7.98		9.52		10.18	10.57			11.90	15.90	
63-125 um	7.86		9.52		10.18	10.57			11.90	15.90	
125-250 um	7.80		9.46		10.16	10.57			11.93	15.90	
ID 354	7.00		5.10		10.10	10.57			11.55	13.50	
0-25 um	8.18	8 34	9 46	9 93	10 18	10 72			12 14	16.01	16 47
25-63 um	7.89	0.54	9.48	5.55	10.18	10.56			11.92	15.01	16.35
63-125 um	7.85		9,39	9.91	10.18	10.54			11.91	15.88	16.49
125-250 um	7.05		9.35	5.51	10.10	10.54			11.71	15.00	16.48
123-230 μm	1.15		5.55		10.10	10.00				13.00	10.40
0-25 um	8.08			9.80					11 01		
25.62 μm	8.00			9.80					11.71		
23-05 μm	8.07			9.00							
125 250 μm	8.00			9.70							
125-250 µm	8.04			9.81							

High-Mg NVP	CF											
ID 355												
0-25 μm	8.08			9.83		10.48		11.	86			
25-63 μm	8.06			9.74		10.50						
63-125 μm	8.04			9.79		10.50						
125-250 μm	8.05			9.79								
ID 356												
0-25 μm	8.08		9.42	9.89	10.23	10.51	10.79	11.43		14.42		
25-63 μm	7.96		9.48	9.89	10.20	10.51		11.30		14.43	15.74	
63-125 μm	7.87		9.42	9.89	10.24	10.49				14.42	15.78	
125-250 μm	7.86		9.39	9.89		10.51	10.79			14.42	15.74	
ID 357												
0-25 μm	8.15	8.38	9.39	9.90	10.23	10.52	10.79	11.44		14.41	15.73	
25-63 µm	7.89		9.42	9.91	10.21	10.53	10.77	11.33		14.42		17.46
63-125 μm	7.83		9.40	9.91	10.22	10.46	10.79	11.34	13.77	14.48	15.63	17.51
125-250 μm	7.81		9.37	9.91	10.21	10.49	10.79	11.34	13.77	14.48	15.70	17.54
ID 181												
0-25 μm	8.08			9.80				11.	91			
25-63 µm	8.07			9.80								
63-125 μm	8.06			9.78								
125-250 μm	8.04			9.81								

Table 4. Band positions of the studied mixtures. In Bold: position of the CF of the respective grain size fractions. (In μm).





Figure 2















- ▲ Bulk △ Micro Synthetic Mercury Glasses (Morlok et al., 2017, 2020a)
- Bulk Synthetic Planetary Glasses (Morlok et al., 2020a)
- Micro
- Impact Rocks and Glasses (Morlok et al., 2016a and b)
- × Glass Basalt Evaporation Experiments (Morlok et al., 2020b)





