Separation of Microplastic Particles from Sewage Sludge Extracts Using Magnetic Seeded Filtration

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ABSTRACT

Microplastic particles (MP) are efficiently retained in wastewater treatment plants and enriched in sewage sludge. For monitoring MP contents in wastewater systems, sewage sludge is thus well suited, but also requires an isolation of MP from the sludge matrix, as other sewage sludge components may interfere with the MP identification and quantification. Although organic matter in sludge samples can be removed through acid and enzymatic digestion procedures, cellulose-mainly from toilet paper-remains in the digests, due to its high chemical resistivity and similar density to MP. We apply the separation concept of magnetic seeded filtration to isolate MP through selective hetero-agglomeration with magnetic seed particles. MP and cellulose differ in their hydrophobic properties and we investigate to what extent these differences can be exploited to selectively form MP-magnetite hetero-agglomerates in the presence of cellulose. These hetero-agglomerates are subsequently separated using a magnet. Five MP types (Polyethylene terephthalate (PET), polypropylene (PP), low density polyethylene (LDPE), polyvinyl chloride (PVC) and polystyrene (PS)) and cellulose particles were mixed in different combinations with both hydrophilic and hydrophobic (silanized) magnetite particles. PET, PP, LDPE and PS only poorly agglomerated with pristine (hydrophilic) magnetite, but efficiently formed hetero-agglomerates with hydrophobic magnetite and were successfully removed from suspensions (80–100%). PVC agglomerated more efficiently with pristine than with hydrophobic magnetite and cellulose only agglomerated to a limited extent with either hydrophilic or hydrophobic magnetite, resulting in a high process selectivity. Results from experiments conducted at different ionic strengths and with hydrophilic and hydrophobic magnetite suggest that the agglomeration process was dominated by hydrophobic interactions. Enzymatic and oxidative treatment of the MP only marginally affected the separation efficiencies and (treated) MP spiked to sewage sludge extracts were successfully recovered using magnetic seeded filtration.

1. Introduction

The superior properties of plastic materials, such as light weight, moldability and durability combined with low-cost production have lead to an enormous increase in the annual production of plastics since the start of the mass production in the late 60-ties and to its accumulation in the environment (Andrady, 2017; Birch et al., 2020; Geyer et al., 2017; Jambeck et al., 2015; MacLeod et al., 2021; Ostle et al., 2019). The ongoing fragmentation of larger plastic items due to mechanical and photochemical weathering results in the formation of microplastic particles (MP) referring to particulate, synthetic polymers with diameters of less than 5 mm (e.g., Hartmann et al., 2019; Thompson et al., 2004), which have been reported from regions all over the globe.

The identification and quantification of MP in various matrices is currently achieved either on an individual particle level or based on total MP quantities in a sample. In the first case, vibrational spectroscopic measurements (e.g., Fourier transform - infrared (FT-IR) or RAMAN) are conducted at spatial resolutions of ≈ 1–10 μm (Anger et al., 2018; Ileva, 2021; Pico et al., 2019; Primpke et al., 2020 and in the second case, mass spectroscopic methods, such as pyrolysis gas chromatography (GC), mass spectrometry (MS) (Dierkes et al., 2019; Funck et al., 2020; Okoffo et al., 2020) or thermal extraction desorption (TED) GC-MS (Duemichen et al., 2019; Eisenraut et al., 2018; Yakovenko et al., 2020) are used. In both cases, however, interference with matrix compounds challenge the analytical techniques and several sample preparation techniques have been developed and tailored for specific matrices (Blaesing and Amelung, 2018; Han et al., 2019; He et al., 2018; Junhao
therefore, received considerable attention and several extraction pro-
cedures tailored for such matrices have been suggested (Hurley et al.,
2018; Sujathan et al., 2017). Philipp et al., 2022 showed that easily
degradable organic materials can be mineralized using a combination of
oxidative and enzymatic digestions and inorganic (sand) particles can be
separated from the MP based on density differences. However, cellulose
materials remain in the digests due to similar size, density and chemical
resistance of cellulose particles and MP.

As density separation and selective (enzymatic) digestions both
failed to separate cellulose from MP, alternative separation methods
have to be considered. Thereby, magnetic seeded filtration (MSF) seems
very promising, which is also supported by initial results published by
Grbic et al. (2019). Magnetic seeded filtration (MSF), also referred to as
magnetic flocculation (Tsouris et al., 1995) and magnetic adsorption
(Franzreb, 2020) is a solid-liquid separation process, which has been
used in various applications (Franzreb, 2020). The approach is based on
the selective hetero-agglomeration between magnetic seed particles and
the (nonmagnetic) target particles. The agglomerates containing the
magnetic seed particles are separated by magnetic separation, whereas
permanent magnets can be used. MSF can also be applied to non-particulate
products, in which case the target analytes are adsorbed to the surface of the magnetic seed particles. This separation process has
for example been applied for harvesting microalgae (Wang et al., 2015),
removing oil leaks (Mirshahghassemi et al., 2017), and purifying pro-
teins (Ebeler et al., 2019; Franzreb et al., 2006). Furthermore, MSF was
applied to remove phosphates (Du et al., 2019) and fine particle matter
(Chin et al., 2006; Gray et al., 1994; Wan et al., 2011) from wastewater
and natural organic matter (NOM) during pre-treatment of drinking
water production on an industrial scale (Drikas et al., 2011). Employing
MSF for the separation of micro- and nanoplastics is currently exten-
sively researched by multiple groups, with ever increasing possibilities
regarding the magnetic seed material (Groppe et al., 2022; Misra et al.,
2020; Shi et al., 2022a). Recent studies show that MSF is capable of
achieving high separation efficiencies for microplastics Rhein et al.
(2019b) under laboratory conditions and identified benefits included
the high material specific selectivity and the applicability in dilute

2. Materials & Methods

2.1. Magnetic seeded filtration theory

The key process, defining the efficiency of MSF is the hetero-
agglomeration between magnetic (and non-magnetic) target particles.
The agglomeration kinetic between two particles $i$ and $j$ resulting in
agglomerate $ij$ can be expressed according to relation 1 (Elimelech,
1998).

$$\frac{dN_{ij}}{dt} = \beta_{ij} N_i N_j$$

$N_i$ and $N_j$ describe the number concentrations of agglomerating par-
ticles. To form an agglomerate, two particles must collide, which is
quantified by the collision frequency $C_{ij}$. In this study, agglomeration is
governed by fluid flow rather than diffusion due to large particle di-
ameters and $C_{ij}$ can therefore be approximated according to Eq. 2 (Chin
et al., 1998; Elimelech, 1998; Han and Lawler, 1992). The collision
frequency is determined by the particle radii and mean shear rate in the
system $G$.

$$\beta_{ij} = \frac{4}{3} (r_i + r_j) G$$

Upon collision, two particles must “stick” together, the probability of
which is expressed through the collision efficiency $\alpha$ in relation 1. The
calculation of $\alpha$ is non-trivial, especially in the shear-controlled regime,
as it requires calculation of particle trajectories under consideration of
the flow conditions (Elimelech, 1998; Han and Lawler, 1992). The relation between $\alpha$ and other relevant parameters is given by Reerink and
Overbeek (1954) and Selomulya et al. (2003).

$$\alpha = \exp \left( \frac{-C_1 \left( 1 - \frac{r_i}{r_j} \right) (r_i r_j)^{1/3}}{C_2} + \exp \left( \frac{-E_{m,a} \alpha_{m,a}}{k_B T} \right) \right)$$

Factor (a) describes the size dependency of $\alpha$ according to Selomulya
et al. (2003) and Snoo et al. (2007) with empirical constants $C_1$ and $C_2$.
For increasing particle size as well as decreasing particle size ratio $r_i/r_j$, the collision efficiency and therefore the probability of an
agglomeration is reduced due to increasing inertia of the particles.
Factor (b) is an approximation for the maximum collision efficiency of
the diffusion-controlled regime (Reerink and Overbeek, 1954) and is
governed by the particle-particle interaction energy $E$. In the traditional
DLVO approach (Derjaguin, 1954), the attractive van der Waals
interactions and (mostly) repulsive electrostatic interactions are considered, whereas this work further requires the inclusion of attractive hydrophobic interactions. For a more detailed description, the reader is referred to Christenson and Claesson (2001), Elimelech (1998), Israelachvili (2011) and to SI.1.

After the hetero-agglomerates are formed, they are separated on the basis of their newly gained magnetic properties. The magnetic force given in Eq. 4 needs to overcome the drag of the hetero-agglomerates. It is primarily dependent on the gradient of the magnetic field strength $\nabla VH$, the magnetization $M$ and partial volume $V_M$ of the magnetic component. It is referred to Svoboda (2004) for a more profound description of magnetic separation and the relevant material properties.

$$F_{Mag} = \mu_0 V_M \nabla VH$$  \hspace{1cm} (4)

After separation, the agglomerates can be further processed to recycle the magnetic seed material and also recover the separated non-magnetic fraction. Agglomerates are either broken up through mechanical stress, or exposed to high temperatures or solvents to extract the non-magnetic component. A previous study (Rhein et al., 2021) showed high recovery rates and low performance losses over the course of multiple separation cycles for three different recycling approaches.

### 2.2. Preparation of MP fragments

Polyethylene terephthalate (PET), polypropylene (PP), low-density polyethylene (LDPE) and polystyrene (PS) beads were purchased from Goodfellow GmbH (Germany) with 5 mm diameter. To generate MP fragments of smaller sizes ($\sim 100 \mu m$), 5 g of the respective beads were ground in a ball mill (M400, Retsch GmbH, Germany) largely following the protocol of Seghers et al. (2021). To cool the 35 mL stainless steel buckets containing both the milling ball with a diameter of 20 mm as well as the MP beads to cryogenic temperatures, they were put in liquid nitrogen for 10 min. The materials were ground several times (PET: 5, PP: 16, LDPE: 15, PS: 5), until the desired size was reached. One grinding cycle consisted of 1 min of milling at $tf = 30 \text{ s}^{-1}$ and 1.5 min of cooling in liquid nitrogen.

### 2.3. Silanization of magnetite

Hydrophobic magnetite was obtained through silanization of magnetite powder (Kremer Pigments, Germany) with hexadecyltrimethoxysilane (HDTMS, > 85% technical grade, Sigma-Aldrich), mainly following the procedures described by Frickel et al. (2010), Grbic et al. (2019) and Ji et al. (2017), which vary in solvent, the amount of added water (if any) and the amount of added HDTMS per surface area of particles. A particle surface specific HDTMS concentration of $4 \times 10^{-4} \text{ g g}^{-1}$ ensures sufficient excess to guarantee complete silanization of the magnetite particles. Assuming perfect spheres and neglecting particle porosity, the required amount of HDTMS can be calculated for different magnetite particle sizes (see SI.2 for more information).

Four of magnetite particles were added to 50 mL plastic tube together with 36 mL of analytical grade ethanol (> 99.9%, EMSURE, Merck, Germany), 4 mL of nanopure water and 0.85 mL of HDTMS. The suspension was mixed in an overhead shaker (Reax 2, Heidolph Instruments, Germany) for 2h. The silanized magnetite particles were separated with a permanent magnet and the remaining liquid was discarded. Following this, the magnetite particles were washed three times with 30 mL of ethanol (technical grade) with magnetic separation of the magnetite particles between each washing step. The washed and silanized particles were dried at 40°C for 17 h and particles from the same batch were used for all of the experiments. Throughout this work, hydrophobic magnetite is abbreviated with Mag-C16, indicating the alkyl chain length of HDTMS.

### 2.4. Treatment of MP

To extract MP particles from complex matrices, including sewage sludge, different protocols haven been suggested, most of them including combinations of enzymatic and oxidative digestion steps (see Section 1). This treatment, however, may induce changes of the surface properties of the MP, which may affect their agglomeration behavior during MSF. To assess these impacts, MP were exposed to enzymatic and oxidative treatment following the sample preparation protocol for sewage sludge described in Philipp et al. (2022). In brief, MP were first exposed to Fenton reagents by adding the MP to a mixture of 10 mL H₂O₂ (35%), 5 mL deionized (DI) water, 1 mL Fe(II)SO₄ ($7 \times H₂O, 2\text{mM}$) and 1 mL protocatechueic acid (2 mM). After a reaction time of 1 h on a horizontal shaker at room temperature and an additional 12 h at 40°C, MP were recovered by filtration and incubated with 50 mL phosphate citrate buffer (50 mM, pH = 5), 0.5 g cellulose (extracted from Aspergillus niger; Sigma-Aldrich, No. 22178) and 10 mg sodium azide at 40°C for 72 h. MP were recovered again by filtration and rinsed with DI water. MP following these treatments are referred to as processed MP in the remainder of the manuscript. Attenuated total reflectance (ATR) - Fourier Transform (FT) - Infrared (IR) spectra recorded on the MP before and after the digestion steps were identical (see Fig. S1.3), demonstrating that the structure of the MP was not affected by the digestion procedure.

### 2.5. Analytical particle solvent extraction

The analytical particle solvent extraction (APSE) was performed following Sygusch and Rudolph (2021) to quantify the wettability of particulate systems. Twenty-fivemg dry particle powder was added to 5 mL of ultrapure water in a 15 mL plastic tube and dispersed in an ultrasound bath for 2 min. Subsequently, 5 mL of 1-Octanol (analytical grade, > 99.9%, Sigma-Aldrich) was added and the mixture was again dispersed for 2 min in an ultrasound bath. Then, the suspension was mixed in an overhead shaker (Reax 2, Heidolph Instruments, Germany) for 20 min. After mixing, the suspension was allowed to rest until a clear phase separation between octanol and water was apparent. Then, 5 mL of the top (octanol) phase were collected with a pipette, and the remaining aqueous fraction was decanted and the tube rinsed with ultrapure water. Both fractions were dried at 40°C and weighted to determine the particle masses in the octanol and in the aqueous phase. The mass fraction in the octanol phase $X_{Octanol}$ was calculated according to Eq. 5 and serves as measure for hydrophobicity: Higher values of $X_{Octanol}$ indicate more hydrophobic particles as they tend to accumulate in the non-polar phase. All extraction experiments were performed in triplicate.

$$X_{Octanol} = \frac{m_{Octanol}}{m_{Octanol} + m_{W_{H₂O}}}$$  \hspace{1cm} (5)

The challenge of this procedure is the accumulation of particles at the liquid-liquid interface. This may either happen directly or due to sedimentation of particles in the octanol phase. The latter was remedied by decreasing resting time, however the APSE did not yield reliable results for MP, as most of the material was found at the liquid-liquid interface. The results of the APSE were used to both validate the silanization of the magnetite particles discussed in Section 2.3 and to characterize the behavior of the cellulose particles.

### 2.6. Particle properties

PET, PP, LDPE and PS particles were prepared through cryo milling as explained in Section 2.2, and polyvinyl chloride (PVC) particles were purchased in powder form. Pulp fibers (J. Rettenmaier & Söhne GmbH, Germany) commonly used as filter aids were used as surrogate for cellulose Bächle et al. (2021). Magnetite particles (Kremer Pigmente, Germany) were used as magnetic seed material and have been described in Rhein et al. (2019b). All particle types were investigated by scanning electron microscopy (SEM) and the resulting images, along with the
operational conditions of the SEM are shown in Fig. SI.2. Besides revealing the fibrous shape of cellulose particles, distinct differences between the cryo-milled particles (PET, PP, LDPE, PS) and PVC are apparent: PVC particles exhibit smooth surfaces and spherical shapes, whereas the cryo milling process led to diverse and often flattened shapes with rugged particle surfaces. The cumulative size distributions of all particle types (MP, magnetite, cellulose), derived from static laser light scattering measurements (LS13 320 XR, Beckman Coulter) are shown in Fig. SI.5. The fraction of all types of MP were within one log10 unit and ranged from 150 μm (PVC) to 450 μm (LDPE/PP). The cumulative size distribution of the cellulose particles was very similar to the cumulative size distributions of the MP, however, this needs to be interpreted carefully due to the fibrous shape of the cellulose particles. The X50% of the magnetite particles was roughly two orders of magnitude smaller than both the MP and cellulose particles resulting in a higher collision probability due to higher number concentrations and a higher volume specific surface area compared to the MP, which enhances the hetero-agglomeration process.

The pristine magnetite particles were hydrophilic, but were made hydrophobic through the silanization process (Table 1). Cellulose fibres were hydrophilic. Strong accumulation of MP at the the liquid-liquid interface during the APSE process hampered an assessment of the hydrophobic properties of the MP and contact angle measurements on polymer films of the same material from the same manufacturer were performed in triplicate to assess the hydrophobicity of the MP. Contact angles were measured on films, as determining the contact angle of hydrophobic particles, subsequently filtered onto membranes (Whatman, 25 mm diameter, 0.2 μm pore size, Sigma-Aldrich) and dried at 40 °C. The relative error of the gravimetric analysis was determined by comparing the mass of the membranes before and after filtration. The experiments were performed in triplicate. The relative error was almost constant over the various concentrations with an average value over all measurements of Δrel = 6.2% ± 6.2% (see SI.7).

2.7. Gravimetric analysis

Gravimetric analyses were performed on an AX205 DeltaRange balance from Mettler Toledo, USA, with an accuracy of 0.01 mg. Prior to the separation experiments, the accuracy of the gravimetric analysis was assessed by spiking different masses ranging from 5 mg to 40 mg of PET to a 50 mL glass bottle. These suspensions were then filtered onto membranes (Whatman, 25 mm diameter, 0.2 μm pore size, Sigma-Aldrich) and dried at 40 °C. Visible light microscopy images of these filters were recorded on a Nikon Eclipse Ni with an automated stage and using a Nikon PlanFluor 4x0.13 WD17.2 objective.

An area of 4500 × 4500 px² corresponding to ≈ 32 × 32 mm² (the largest square that can be fitted into the 47 mm filter) was selected for image analysis. The cellulose fibres in the images of the separated fractions were manually colored and then segmented to quantify the cellulose fibres based on binarized images. Cellulose was clearly distinguishable from the MP due to their characteristic, fibrous shape and the slightly different color. The binary images were processed using

![Image](image-url)

### Table 1

Properties of the particulate materials and of the polymer films (PET: Polyethylene terephthalate; PP: polypropylene; LDPE: low-density polyethylene; PVC: polyvinyl chloride, PS: polystyrene).

| Material | Magnetic (M) | Non-magnetic (NM) | PET | PP | LDPE | PVC | PS |
|----------|--------------|-------------------|-----|----|------|-----|----|
| Manufacturer | Kremer Pigments | J. Rettenmaier & Sohne GmbH | Goodfellow | Goodfellow | Goodfellow | Goodfellow | Goodfellow |
| Density [kg/m³] | 5200 | 1500 | 1400 | 900 | 920 | 1400 | 1100 |
| X₁₀₂₅ [μm] | 2.1 | 110 | 150 | 230 | 210 | 110 | 130 |
| X₅₀ [μm] | 7.2 | 310 | 310 | 450 | 450 | 150 | 280 |
| X₉₅ [μm] | 21 | 590 | 490 | 650 | 1100 | 210 | 490 |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Contact angle (θ) | | | | | |
| Fraction Octanol X₀ [%] | 6.5 ± 3.0 | 12 ± 6.6 | | | | | |
| Fraction Octanol X₀ [%] | 90 ± 4.5 | 6.5 ± 3.0 | | | | | |

* Technical data sheet manufacturer | ^ Measured with LS13 320 XR laser diffraction particle size distribution analyzer from Beckman Coulter Inc, USA. See Fig. SI.5 for full distribution | † No comparable measurement for particle powders | © Measured with the Dataphysics OCA-20 contact angle analyzer. Static contact angle measured by sessile drop method at 20 °C and 70% relative humidity with films of the same material by the same manufacturer. Ultrapure water with drop volume 10 μL | ‡ No reliable results due to accumulation in the liquid-liquid interface. All values are rounded to 2 significant digits
the open source software Fiji Schindelin et al. (2012) to determine the total 2D area of the cellulose fibers. As reference, the same initial amount of cellulose fibers as was used in the 3-component experiments (see Table 2), was filtered and processed as described above. As these samples contained cellulose fibers only, the identification of the cellulose fibers on the recorded images was automated using specific thresholds for segmenting the images. The separation efficiency of cellulose fibers was estimated by comparing the total area of cellulose fiber on the filters resulting from separated samples to the respective area from the cellulose reference filters. The workflow of the image analysis pipeline is schematically shown in Fig. S1.7.

2.9. Experimental design

The experimental design is depicted in Fig. 1, with the experimental parameters listed in Table 2. Additional information about the absolute masses is given in Table. SI.2. Desired amounts of MP and/or cellulose particles were weighted into a50 ml glass bottle (Schott AG,Germany) and magnetic particles into a separate weighting bottle. Then, ultrapure water (atrum pro, Sartorius, France) and a4 M NaCl solution (analytical grade, EMSURE, Merck, Germany) were added to reach the liquid volume of40 ml and the ionic strength (I) of the given experiment. The pH was 7.05 (I = 0.01 M, T = 21 °C). The magnetic particles were added to the MP suspension and the weighting bottle was rinsed with5ml of ultrapure water. Agglomeration between magnetic and non-magnetic particles was induced by putting the glass bottle for10 min on a horizontal shaker (KS260 basic, IKA, Germany) at250 rpm. Subsequently, the bottle was removed and placed next to a permanent magnet for2 min to separate the magnetic agglomerates from the non-magnetic particles. The supernatant was carefully decanted while holding the permanent magnet in place.40 ml of an NaCl solution with the same ionic strength as the respective experimental sample was added to rinse the glass bottle. A second magnetic separation was performed analogously but with a reduced magnetic separation time of1 min. The collected supernatants were filtered onto a previously weighted membrane filter (Whatman,25 mm diameter,0.2 μm pore size, Sigma-Aldrich) and dried at40 °C until constant weight, but at least for12 h. In the 3-component experiments (MP, cellulose and magnetic particles), the separated fraction was resuspended, filtered and also weighted. This allowed the quantification of particle losses required for the calculation of the separation efficiencies. As this study was focused on the agglomeration behavior of the MP in the context of MP analysis, the possibilities to quantify the magnetic seed particles and ultimately the economy of the process were less relevant and thus not addressed in this work.

The separation efficiency for non-magnetic particles in the 2-component system (T_{NM,2-comp}) was calculated according to Eq. 6. The assumption was that all magnetic particles added to the system (m_{M,0}) were separated and therefore, particles in the supernatant (m_{NM}) corresponded to not separated non-magnetic material at the end of the experiment (m_{NM,E} = m_{NM}) which was ensured by initial experiments.

\[ T_{NM,2-comp} = 100\% \left(1 - \frac{m_{NM,E}}{m_{NM,0}}\right) = 100\% \left(1 - \frac{m_{NM}}{m_{NM,0}}\right) \]  

During 3-component experiments, the separated fraction was recovered and weighted to estimate particle losses during the procedure.

The initial mass of non-magnetic particles added to the system is denoted by m_{NM,0}.

The mass loss\Delta m over the individual 3-component experiments was calculated as the difference between the total mass of the particles recovered (mass of the supernatant (m_{NM}) and of the separated fraction (m_{comp})) and the total mass of the added particles at the beginning of the experiment (m_{MP,0}, cellu\]lose (m_{Cellulose,0}), magnetite (m_{magnetite,0}). Assuming that the mass losses can be equally distributed between the supernatant (SN) and separated (SEP) fraction and that cellulose is separated to the same degree as observed in the 2-component system (T_{Cellulose,3-comp} = T_{Cellulose,2-comp})

The separation efficiency of the MP in the 3-component system (T_{MP,3-comp}) was calculated according to Eq. 7. The validity of this assumption was assessed through microscopic image analysis of the separated fractions and is covered in Section 3.2.

\[ T_{MP,3-comp} = 100\% \left(1 - \frac{m_{NM} - \Delta m}{m_{NM,0}} - \frac{m_{Cellulose,0}}{m_{MP,0}} \left(1 - \frac{T_{Cellulose,3-comp}}{100}\right)\right) \]  

3. Results & Discussion

3.1. 2-Component System

The separation efficiencies when using untreated, hydrophilic magnetite (Mag) were around10% for PET, PP, LDPE and PS (Fig. 2) and cellulose was separated to>20%. However, separation efficiencies exceeding80% were obtained for PVC. The separation behavior drastically changed, when using hydrophobic magnetic particles (Mag-C16). PP, LDPE and PS were almost completely separated and separation efficiencies of>80% were obtained for PET. In contrast, the separation efficiencies of PVC and also for cellulose significantly decreased and were around30 – 40% for PVC and10% for cellulose.

These trends correlated with the hydrophobic properties of the respective polymer films listed in Table 1, except for PVC which is discussed separately. PP, LDPE and PS exhibit contact angles>90° and can be considered as hydrophobic (Israelachvili, 2011). The hydrophobic behavior of these polymers and Mag-C16 results in a strong attractive interaction between these particle types, enabling their agglomeration and subsequent magnetic separation. The origin of the hydrophobic force on a molecular level can be retraced to the rearrangement of water molecules between two hydrophobic surfaces (Israelachvili and Pashley, 1982), but hydrophobic particle-particle interactions are acting on a much larger length scale and are currently intensively debated. There seems to be a consensus that the origin of the long ranged attractive force is due to the bridging of nanobubbles attached on hydrophobic surfaces (Köckich, 2019), but Iishida et al. and Ishida et al. (2018) demonstrated that even in the absence of such bubbles, strong attractive forces can be measured. When hydrophilic magnetic particles are used instead, the formation of hetero-aggregates is suppressed, which is reflected by the low separation efficiencies of the polymers. The separation efficiency for PET in combination with Mag-C16 was around80% and thus lower than the corresponding efficiencies for the PP, LDPE and PS. The slightly lower separation efficiency for PET may be explained by the lower contact angle of around77° resulting in a less efficient formation of hetero-aggregates.

The APSE results in Table 1 show that cellulose accumulates in the aqueous phase and can, thus, be considered as hydrophilic. The lower separation efficiencies when using Mag-C16 instead of hydrophilic magnetite can be explained by the formation of Mag-C16 homo-
This reduces the total number of magnetic particles available to form hetero-agglomerates and reduces the aggregation kinetics (Eq. 1). This effect applies to all MP types, however, is offset for PET, PP, LDPE and PS by the additional attractive hydrophobic interaction leading to an increased collision efficiency $\alpha$.

Increasing the ionic strength by a factor of 100 from 0.01 M to 1 M did not significantly change the separation efficiency (Fig. 2), which is in contrast to previous studies on MSF (Rhein et al., 2019a, 2020, 2019b). At pH 7 all investigated particle systems are expected to be negatively charged (see Section 2.6), resulting in an electrostatic repulsion. According to SI.1 an increase in ionic strength reduces the range of the repulsive electrostatic interactions $\kappa^{-1}$1, lowers the energy barrier $E_{\text{min}}$ and leads to an increased collision efficiency $\alpha$ in Eq. 1. This facilitates the formation of aggregates and ultimately leads to an increase in the separation efficiency. The independence of the separation efficiencies from the ionic strength, therefore, suggests that the compression of the electric double layer did not change the aggregation behavior and that therefore, electrostatic forces were not controlling the hetero-aggregation process. Based on the results from previous studies (Rhein et al., 2019a, 2020, 2019b), only a negligible repulsion is expected at $t = 1$ M. The fact that this did not lead to an increase in the separation efficiency (in the experiments conducted with pristine magnetite), therefore indicates that the van der Waals attraction between particles is too weak to induce an agglomeration.

The most plausible explanation for this phenomenon is found in the particle diameters, which are orders of magnitude larger than in Rhein et al. (2019a, 2020, 2019b): Whereas the van der Waals interaction increases proportionally to the particle radius, inertial forces or flow effects increase proportionally to the volume of the particle, i.e. $r^3$. A factor of 10 in diameter, consequently increases the ratio of inertial to van der Waals forces by a factor of 100. This effect is accounted for by the factor $a$ in Eq. 3 and drastically reduces the collision efficiency for increasing particle diameter. Furthermore, the (cryo)-milling process likely increased the surface roughness of the polymers (see Fig. SL2), which may lower their effective Hamaker constants by several orders of magnitude (Valmacco et al., 2016). The reduced van der Waals interactions thus explain the low separation efficiencies observed for PET, PP, LDPE and PS in combination with pristine magnetite. Similarly, cellulose exhibits diverse, fibrous shapes with rough surfaces, explaining the low van der Waals attraction and separation efficiency. Although hydrophobic interactions also increase proportionally to the particle radius (Christenson and Claesson, 2001), they nevertheless are able to induce an agglomeration despite the greatly increased inertial effects.

Multiple studies (Christenson and Claesson, 2001; Rabinovich and Derjaguin, 1989) have shown that depending on the material system, the hydrophobic behavior can outperform van der Waals interactions by several orders of magnitudes and are therefore in agreement with the increased separation efficiencies of PET, PP, LDPE and PS when agglomerated with Mag-C16 as shown in Fig. 2.

The separation behavior of PVC was different to the other MP and substantially decreased from $\approx$ 80% for pristine (hydrophilic) magnetite to $\approx$ 35% for Mag-C16. This behavior is similar to the behavior observed for cellulose and generally indicates hydrophilic surface properties. Mag-C16 particles most likely homo-agglomerated during the experiment with PVC, which reduced the agglomeration kinetic (Eq. 1). The hydrophobic behavior of the PVC particles, however, is in apparent discrepancy to the results from the contact angle measurements of $\approx$ 86° conducted on PVC films (Table 1), suggesting hydrophobic surfaces. Energy dispersive x-ray analyses conducted in the SEM (SEM-EDX) of a PVC film and of selected PVC particles revealed significantly different chlorine (Cl) contents (see SL.5), which indicate different surface properties of the particles compared to the films. Thus, the contact angle determined for the PVC films may not be transferable to the PVC particles. Based on the agglomeration and separation behavior of the PVC particles we predict a lower contact angle of the PVC particles compared to the PVC films, consistent with the hydrophilic behavior of the PVC particles. Due to the experimental challenges outlined in Section 2.5, however, we were not able to experimentally assess the contact angle of the PVC particles. In addition to the surface properties, also the surface roughness of the PVC particles may have contributed to their hydrophilic behavior during hetero-agglomeration. In contrast to other MP, which were ground in a ball mill under cryogenic conditions, PVC was directly obtained in powder form and with a smaller particle size compared to the other MP (see Table 1 or Fig. SL2). The high separation efficiencies for PVC of $\approx$ 80% when pristine (hydrophilic) magnetic particles were used, suggest that van der Waals interactions exceeded inertial forces and were inducing the formation of agglomerates. This can be attributed to both the smaller particle size, reducing inertial forces, and larger effective Hamaker constants resulting from a smoother particle surface due to the manufacturing process of the PVC particles (see Fig. SL2). Additionally, the experiments with PVC showed larger standard deviations, which may indicate weaker (van der Waals) contact forces in the agglomerates. Small variations in the experimental procedure may therefore result in agglomerate breakage and explain the variance in separation efficiency. Although it is difficult to quantify, differences in particle shape (see Fig. SL2) may also influence the particle-particle collision and therefore the separation efficiency.

All processed MP types, with the exception of PET, behaved very
similar to the pristine ones during MSF (Fig. 3). Processed PP, LDPE and PS were almost quantitatively separated and processed cellulose was hardly separated at all. The average separation efficiency of PVC slightly increased to 40–50%, but due to the large standard deviations, the effect is not considered significant. Table 1 further supports this, as the contact angle of PVC films remained unchanged through processing. The separation efficiency for processed PET, however, was reduced to 20–30%. This reduction in separation efficiency can be explained by chemical changes modifying the surface properties of the PET which was reflected in a decreased contact angle of the processed PET of around 65° (Table 1). The exposure of PET to strong oxidation agents (Fenton reagents) and to specific enzymes likely caused the formation of carboxylic acid end groups (Gewert et al., 2015) rendering the surfaces more hydrophilic and explaining the change in the contact angle. The oxidation of PET surfaces is known to reduce the hydrophobicity i.e. increase the wettability and is a desired effect in designing membrane materials (Pivaró et al., 2007; Gotoh et al., 2018; Korolkov et al., 2015).

Although the contact angles of processed PP and PS were also reduced compared to the pristine materials, the separation efficiency for these polymers remained constant at close to 100%. Thus, with the exception of PET, the separation efficiency of the pristine and the processed MP were similar, making the MSF a promising approach to isolate MP from digests of complex matrices (e.g. sewage sludge). Even for processed PET, the difference in the separation efficiency compared to cellulose is significant indicating a selective separation. By increasing e.g. agglomeration time or performing a multistage separation, the degree of separation for PET might be increased whereas cellulose is expected to remain in the suspension.

In summary, the results in Figs. 2 and 3 show that the separation efficiency is dominated by hydrophobic and van der Waals interactions. The ionic strength did not influence the separation efficiency, indicating that the electrostatic interactions were negligible for the investigated systems. Separation of hydrophobic nonmagnetic particles can be realized by using hydrophobic magnetic particles without increasing the separation efficiency of hydrophilic cellulose. Whereas the separation efficiencies for PP, LDPE and PS were mostly below 10% for hydrophilic magnetic particles, an almost complete separation was achieved with hydrophobic magnetite, independent of a previous processing step. The different behavior of PVC suggests that the surface roughness plays a key role in MSF. Rough surfaces either created during the cryo milling process in the case of PET, PP, LDPE and PS or inherently present in the case of fibrous cellulose particles reduce the effective Hamaker constants which may help to explain the different separation behavior of these particles compared to PVC.

### 3.2. 3-Component System

The results from the 2-component system suggested that a separation between hydrophobic MP and hydrophilic cellulose is possible. However, potential interactions between cellulose and MP may lead to an increase in the separation of cellulose and, thus, a reduced selectivity. A detailed discussion about selectivity in multi-component agglomerating systems is given in (Rhein et al., 2020). In the 3-component system, cellulose and MP can be present in the separated fractions. Therefore, more detailed information about the composition of these fractions in addition to gravimetric analyses were required to assess the separation efficiencies of the MP. In this study, the separated fractions of cellulose in each of the 3-component experiments were quantified by image analysis techniques as described in Section 2.8 and the obtained separation efficiencies of cellulose are shown in Table 3. Original and processed images are given in Fig S1.8.

The separation efficiencies for cellulose in all 3-component systems derived using image analysis tools, were in the range of 0.37%–2.3%, indicating only a limited interaction between cellulose and the MP resulting in cellulose separation efficiencies similar to the 2-component system. Overlapping cellulose fibers, more prominent in the reference sample due to the higher cellulose concentration and manual identification of cellulose fibers in separated fractions both contributed to the uncertainty associated with the determination of separation efficiency in the 3-component system. We accounted for these effects by varying the image processing parameters (brightness/contrast settings for cellulose identification). As the separation efficiencies for cellulose obtained through image analysis were in good agreement with separation efficiencies of cellulose obtained from the 2-component systems, the separation efficiencies for cellulose in the 3-component system were set equal to the respective separation efficiencies determined in the 2-component system ($T_{\text{Cellulose,3-comp}} = T_{\text{Cellulose,2-comp}}$). The separation efficiencies of the MP were, thus, calculated according to Eq. 7 and are shown alongside the 2-component results in Fig. 4 for pristine and Fig. 5 for processed MP. The experiments were performed in suspensions with an ionic strength of 0.01 M and Mag-C16 particles. The cellulose separation efficiency is displayed with corresponding standard deviation as a horizontal line.

Within the standard deviations derived from three replicates, the pristine and processed MP showed an almost identical separation behavior in the 2- and in the 3-component system, which is consistent with the negligible interactions between cellulose fibers and MP. The results demonstrate that a selective separation between MP and cellulose with MSF is possible, and can reach high levels of selectivity. PP, LDPE and PS were quantitatively separated from the liquids, whereas cellulose was only separated to a limited extent. It is remarkable that the results from experiments conducted with PVC consistently show larger standard deviations compared to the results obtained from experiments conducted with the other MP. This may point towards the formation of weak hetero-agglomerates in the secondary minimum, which may be ruptured to various extents during the magnetic separation process. The separation efficiencies for PET and PVC were around 70% and 40%, however, the selectivity of the process was still high. This means that PET and PVC will still be present in the supernatants, but the separated fraction will be almost cellulose-free. The overall separation might, therefore, be increased by extended agglomeration times or a multistage separation, resulting in higher recoveries in real-world samples, should this be necessary.

### 3.3. Separation of polymers from sewage sludge extracts

To assess the potential of the MSF approach to separate MP from digested sewage sludge extracts, additional separation experiments were performed. Digested sewage sludge samples were collected from a full-scale wastewater treatment plant (ARA Neugut, Duebendorf, Switzerland) and processed following the extraction protocol published by Philipp et al. (2022) and described in Section 2.4. The sludge extracts (ESX), essentially consisting of cellulose and polymers already present in the digested sewage sludge are considered as a separate type of nonmagnetic particle system. An initial separation experiment was performed with ESX yielding a separation efficiency of $T_{\text{ESX}} = 16.65\%$, meaning that about 17 wt% of the particulate material present in the ESX was found in the separated fraction following the MSF approach and 83 wt% remained in the supernatant. Subsequently, processed PET

### Table 3

| MP type | PET | PP | LDPE | PVC | PS |
|---------|-----|----|------|-----|----|
| Cellulose 1 | 0.85 | 0.38 | 0.58 | 1.2 | 0.37 |
| Cellulose 2 | 1.6 | 0.72 | 1.1 | 2.3 | 0.69 |
Fig. 2. Separation efficiencies for pristine non-magnetic particle systems (NM: cellulose and different microplastic particle types) in the 2-component system. Green bars refer to experiments conducted with untreated magnetite (Mag) and blue bars refer to experiments conducted with hydrophobic magnetite (Mag-C16). The ionic strength was increased from 0.01 M (left bars, light colors) to 1 M (right bars, dark colors). PET: Polyethylene terephthalate, PP: polypropylene, LDPE: low-density polyethylene, PVC: polyvinyl chloride, PS: polystyrene.

Fig. 3. Separation efficiencies for pristine and processed non-magnetic particle systems (NM: cellulose and different microplastic particle types) in the 2-component system. All experiments were conducted with hydrophobic magnetite (Mag-C16). Blue bars refer to experiments conducted with pristine NM and red bars refer to experiments conducted with processed NM. The ionic strength was increased from 0.01 M (left bars, light colors) to 1 M (right bars, dark colors). Abbreviations for the different microplastic particle types are given in Fig. 2.

Fig. 4. Separation efficiencies for different microplastic particle (MP) types in the presence of cellulose (3-component system). Experiments were conducted with pristine cellulose, pristine MP and hydrophobic magnetite (Mag-C16) (blue bars). The separation efficiencies from the respective experiments in the 2-component system (pristine MP and Mag-C16) are added for comparison (grey bars, see Fig. 3). The ionic strength was 0.01 M and the separation efficiency for cellulose was assumed to be the same as in the 2-component system. Abbreviations for the different MP types are given in Fig. 2.
(3.43 mg ± 0.19 mg), PP (2.34 mg ± 0.16 mg) and PVC (3.30 mg ± 0.03 mg) were spiked to EXS (6.99 mg ± 0.04 mg), resulting in identical volume concentrations of all MP and in similar volume concentrations of the sum of all MP and EXS. After the MSF of these experimental suspensions, both the absolute mass of particles in the supernatant and in the separated fraction were determined gravimetrically. Assuming that the separation efficiencies determined for the 2- and the 3-component systems are also valid for multicomponent systems, the predicted mass of the supernatant \( m_{\text{SN,exp}} \) can be calculated according to Eq. 8

\[
m_{\text{SN,exp}} = \sum_{n=1}^{N} m_{n,0} \left(1 - \frac{T_n}{100}\right),
\]

with \( m_{n,0} \) referring to the initial masses of the individual, spiked MP and solids already present in the sludge extracts (EXS) and \( T_n \) representing the separation efficiencies of the respective non-magnetic components (MP, EXS).

The mass of the particulate material in the supernatant, most likely dominated by the sum of spiked MP and MP/cellulose that were already in the sludge extracts, calculated by 8 was in good agreement with the experimental data shown in Table 4. Thus, the results suggest that MSF can be applied possibly as a final polishing step as part of a sample preparation protocol for the selective separation of MP from complex, environmental samples.

4. Conclusions

Cellulose fibers originating for example from disintegrating toilet papers, are abundant in digested sewage sludge. Recently published MP extraction protocols can efficiently isolate MP from most of the sludge components, but cannot separate MP from cellulose as MP and cellulose have similar densities and are both resistant against chemical and enzymatic digestions. For analytical purposes, e.g. pyrolysis GCMS measurements, the separation of cellulose materials from MP, however, would be highly welcome to avoid possible interference between pyrolysis products from MP and cellulose. This study demonstrated the potential of MSF using hydrophobic magnetic particles as seeding agents to separate MP from cellulose materials. For PP, LDPE and PS an almost complete separation of the MP from the liquids was achieved, whereas cellulose was only found in negligible amounts in the separated fractions. Although, PET and PVC were separated less efficiently from the liquids (≈ 75% and ≈ 40%, see Fig. 4), the MSF was very specific as cellulose mostly remained in the supernatants. Therefore, the respective MP may be enriched in the separated fractions by repeated MSF or, alternatively, the obtained MP fractions may be corrected according to the respective separation efficiencies. The presented MSF approach using hydrophobic seed particles, represents a straight-forward and promising purification step, likely facilitating the quantification of MP extracted from complex matrices. The contrasting results obtained for PVC compared to other MP types emphasize the complexity of real-world MP systems and challenge the use of experimental results obtained from selected / or specifically synthesized MP to assess the fate and behavior of MP in the environment.

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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Supplementary materials

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