



Cite this: *Phys. Chem. Chem. Phys.*,
2026, 28, 790

Predicting polyacrylate–microplastic interactions with atomistic simulation

Timothy M. E. Jugovic,^a Henry E. Thurber,^{ib} Michael T. Robo,^a Woojung Ji,^a
Madeline E. Clough,^{ib} Anne J. McNeil^{ib} and Paul M. Zimmerman^{ib}*^a

Adhesive-coated substrates have been used to non-selectively capture microplastics (MPs) in water. This work explores whether selective MP capture is plausible by modifying the adhesive structure. To find the degree of selectivity, adhesive strength might be estimated by examining surface interactions between specific adhesives and MPs. Herein, we describe the use of atomistic simulations to predict the aqueous adhesive strength via the aqueous work-of-adhesion ($WoA_{(aq)}$), a thermodynamic measure of adhesion. Simulation of four common MPs and five different polyacrylate adhesives were conducted. The simulations show that fluorinated sidechains, due to their low interfacial energies and higher surface energies, exhibit increased selectivity toward polystyrene over other plastics. To provide experimental support for these predictions, probe-tack studies of aqueous adhesion were performed and found to agree with the simulations. Simulations also revealed non-intuitive interactions that govern $WoA_{(aq)}$, arising from the complex intra- and intermolecular interactions that occur when polyacrylates interface with water and MPs. This expanded understanding of the microscopic features of adhesion can be used to design next-generation adhesives and MP remediation technology.

Received 19th September 2025,
Accepted 27th November 2025

DOI: 10.1039/d5cp03631c

rsc.li/pccp

Introduction

Acrylate polymers are versatile materials used in industrial and commercial applications.¹ Products include Plexiglas (poly(methyl methacrylate)), water-soluble thickeners (sodium polyacrylate) and adhesives (poly(cyanoacrylate) and poly(2-ethylhexyl acrylate)).^{2,3} These applications are possible due to the acrylate functionality, which contains an ester-linked side-chain, permitting modular tuning of the polymer properties, including polarity, mass, and volume. As an example, polymers with mixed side-chains like poly(methyl/*t*-butyl acrylate) have higher work of adhesion (WoA), a measure of the adhesive strength, than either homopolymer.⁴ Adhesive applications range from mild pressure-sensitive adhesives (PSAs) such as poly(2-ethylhexyl acrylate) with a $WoA \sim 20$ – 30 mN m^{-1} , to strong cyanoacrylate adhesives with a $WoA > 100 \text{ mN m}^{-1}$.^{1,5} Similarly, the aqueous work of adhesion ($WoA_{(aq)}$), or the strength of an adhesive in water, is important for applications in water and has previously been determined for an epoxy adhesive/carbon reinforced fiber composite interface at $\sim 22 \text{ mN m}^{-1}$.⁶

Microplastics (MPs) are a pervasive form of plastic pollution and are difficult to remediate due to their small size and

heterogenous composition.^{7–14} Moreover, environmental degradation of MPs into even smaller nanoplastics is a growing concern.¹⁵ Capturing microplastics before they are released into the environment is the best option. Once captured, those MP could be recycled into new plastic products. However, the physical separation of different types of microplastics would be challenging given their small size.^{16–18} Slight differences in MP properties might allow for adhesives to selectively bind them, providing a means to simultaneously capture and separate MPs by type. Acrylate polymers are potential materials for this purpose because their adhesive strength can be tailored with structural modifications. A common way to predict adhesion is to compare surface tensions. The surface tension represents the energy of a given surface when in contact with another material (*i.e.*, liquid). Adhesion occurs between two materials when the surface tension is reduced. For plastics, surface tension (γ) spans from non-polar, at $\sim 15 \text{ mN m}^{-1}$, to polar, $\sim 40 \text{ mN m}^{-1}$,^{19,20} where common MP materials (polyethylene, polystyrene, polyamides, and polyesters²¹) are in the range of 15 – 30 mN m^{-1} .²² Unfortunately, the small range in γ mean plastic surfaces are more alike in adhesive properties than they are different.²³ For context, the γ of water is 72 mN m^{-1} ,²⁴ and common metal surfaces have γ of 500 – 1000 mN m^{-1} .²⁵ Selectivity in adhering to less distinct surfaces therefore requires exploiting slight differences in composition that might bias intermolecular interactions, such as π -stacking or H-bonding, between the adhesive and MPs. These intermolecular interactions can also lead to deformation of the

^a Department of Chemistry, University of Michigan, 930 North University Avenue, Ann Arbor, Michigan, 48109-1055, USA. E-mail: paulzim@umich.edu

^b Macromolecular Science and Engineering Program, University of Michigan, 2800 Plymouth Road, Ann Arbor, Michigan, 48109-2800, USA



adhesive-MP interface, leading to changes in polymer conformation and further contributing to changes in adhesive properties. Examining the precise polyacrylate compositions necessary to achieve such selectivity is a significant undertaking. By developing models that are better able to predict adhesive behavior, this process could be accelerated.

The relationship that determines adhesion between surfaces²⁶ can be expressed *via* the difference between their individual surface tensions and their combined interfacial tension, which gives the following equation:

$$\text{WoA}_{(\text{aq})} = \gamma_{\text{adh-H}_2\text{O}} + \gamma_{\text{plas-H}_2\text{O}} - \gamma_{\text{adh-plas}} \quad (1)$$

(adhesive (adh), solvent (H₂O), microplastic (plas); see SI, Section 1 for derivation). The resulting $\text{WoA}_{(\text{aq})}$ is the energy required to separate two surfaces per unit area. The total energy required for separation therefore depends on WoA and the total contact area. The γ for polyacrylates or other surfaces can be usefully conceived as arising from two components: polar and dispersive. Current understanding of γ hold that favorable interaction occurs when surfaces have similar ratios of polar and dispersive γ components. This understanding, while valuable, is not the full story due to emergent interactions in more complex materials.^{27–33} For example, the γ for an adhesive that combines both polar and non-polar co-monomers would be harder to predict. More comprehensive models of γ would allow for robust predictions of relative adhesive ability.

We hypothesized that polyacrylate compositions can be tailored to enable selectivity in binding specific MPs. Acrylate polymers can contain a range of side chains, from non-polar alkyl and fluorocarbon groups to highly polar carboxylate groups. This range provides a large design space. Achieving selectivity would require tuning side chain characteristics, such as polarity, volume, and mass, all of which play a role in determining polymer properties.³⁴ In addition, the polymer length can also affect the γ , as longer polymers have different cohesive behavior.³⁵ Similarly, emergent properties are more difficult to predict. Nevertheless, these interactions may become predictable through computational modeling efforts, as described below.

Computational methods, such as atomistic molecular dynamics (MD) simulations, can be used to build predictive models of polymer structure, behavior, and properties. The most common use of MD simulation is analysis of structure and function of proteins and peptides.^{36–38} This focus has led to most MD forcefields being optimized for amino acids, and other biologically relevant species. Commonly modeled protein interactions, such as quaternary structures, are analogous to adhesion between polymers and microplastics.³⁹ Also, MD forcefields have been constructed to simulate interfacial adhesion^{40,41} as well as behavior and properties of plastic polymers.^{42–49} Recent work on polyacrylate adhesives highlights the power of MD simulation to elucidate properties that are normally difficult to determine using experimental methods.⁵ These previous works inform how interactions between polymers result in unexpected behaviors. Conceptual understanding of how these behaviors arise will enable better predictive models of novel polyacrylate adhesives. To develop design principles for polyacrylates with selective plastic

adhesive properties, we examine structure–property relationships that connect polymer composition with WoA to MPs. The models are comprised of pairs of polymer chains that interact *via* surface contacts, allowing prediction of the microscopic interactions that lead to adhesion at the bulk level. To ensure the reliability of these atomistic models, tests against known literature quantities were conducted and then applied to determine behaviors that govern adhesion. These models provide qualitatively useful concepts for tailoring interactions between acrylates and common MP types, including non-obvious behaviors for fluorine-containing acrylates.

Methods

Polymer generation

Novel polyacrylate compositions were generated using an in-house polymer generation code (see SI for further details). Each polyacrylate strand was constructed from a racemic mixture of 2-ethylhexyl acrylate side chains and a different side chain (R^*) with a composition indicated by its % R^* value (see Chart 1). Each generated polymer strand was relaxed *in vacuo* to a final simulation structure in 1 ns CHARMM MD simulations. For MP surfaces, CHARMM-GUI was used to generate polymer strands of PA-6, HDPE, and PS of sufficient size to approximate macroscopic properties, which were annealed into MP particles of around 10–100 nm³ in size in 1 ns CHARMM-MD simulations (see SI, Section 2 for more details).⁵⁰ These surfaces are representative of pristine MP models, lacking *in situ* changes or modifications that may occur in the environment.

Force field validation

The CHARMM general force field version 4.0 (cgenff) was used to model the interactions, as prior work had parameterized force fields for similar atom types.^{42–49} The forcefields were tested by comparison to known literature values, in particular the density and aqueous interfacial surface tension (γ_{SL}). Table S2 shows the computed and literature aqueous interfacial surface tension (γ_{SL} with SL for solid–liquid) between water and surfaces. Literature values are derived from contact angle (θ_{SL}) and surface energy (γ_{SG} with SG for solid–gas) using the Young–Dupré equation.²⁸ For all examined MPs, γ_{SL} are within 1–2 standard errors of literature or experimental values.^{51–55} The values of γ_{SL} are not available for the novel polyacrylates, so the estimated density is used as an alternative comparison (Table S1). Literature

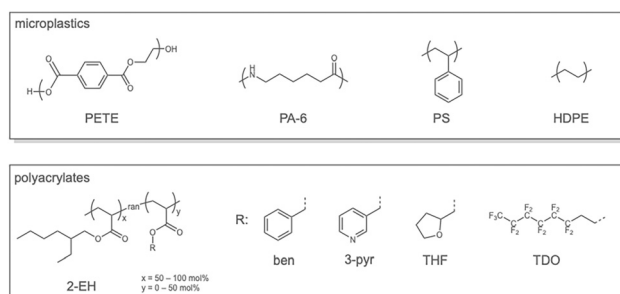


Chart 1 Structures of tested MPs and polyacrylates.



values for density of $R:R^*$ mixed polyacrylates were estimated using the weighted average of the known pure values. Similar agreement is found between the computed and estimated experimental density. Because density is intrinsically tied to the degree of crystallinity, this agreement (Table S1) suggests the MP models are reasonable. The agreement between these values and experiment, combined with the initial parameter validation done for cgenff force field for these bond types, give a measure of confidence in the reliability of our computational results.

Statistical validation

An important concern when analyzing mixed polyacrylates is to sample multiple polymers of the same composition to arrive at meaningful averages for their properties. Another possible statistical issue is clustering of R^* groups within the polymer, resulting in structures that resemble block copolymers more closely than random copolymers. To avoid these issues in simulations, six versions of each % R^* composition were independently generated using an algorithm that randomly distributed the desired % R^* sidechains throughout the structure. These structures are kept at a fixed molecular mass, and polymeric length was varied as composition varied, around 150–300 monomeric units. Each of these six chains was then independently equilibrated for 250 ns for the aqueous and MP boxes. After equilibration, trajectories were simulated for 5 ns with a frame taken every 200 ps to determine an average γ value for each case. This resulted in 60 simulations for each % composition that were then averaged to determine the $WoA_{(aq)}$. Error was quantified as the standard deviation of the combined simulation data for each data point. Outliers were identified using the Q -test and excluded from the reported results. Further description of methods can be found in the SI.

Results and discussion

Our discussion is delineated in four general areas. First, the choice of specific polyacrylates and microplastics to represent realizable copolymers and environmentally relevant plastic types is outlined. Next, the WoA is determined from MD simulations for each microplastic across the span of copolymer compositions. Then, the same atomistic models are examined to determine the acrylate properties that affect adhesive ability for varying polymeric composition. Finally, adhesive simulations are analyzed for behaviors or trends that could be used to separate MPs by plastic type using adhesion.

Chemical space: MPs and acrylates

Four types of plastic were chosen to represent some of the most abundant microplastics in the environment (Chart 1).⁸ In particular, high-density polyethylene (HDPE), polystyrene (PS), nylon-6 (PA-6), and polyethylene terephthalate (PETE). These plastics also span the polarity space of common plastics from purely nonpolar (HDPE) to moderately polar (PA-6, PS). Here, polarity refers to the fraction (in %) of surface energy attributed to polar components (*e.g.*, dipoles) compared to the total energy, which also includes dispersive components.³⁴ The reported

Table 1 Percent polar composition of adhesives and microplastics derived from literature results

Polymer	γ_d (mN m ⁻¹)	γ_p (mN m ⁻¹)	% Polarity
Adhesives			
2-EH	27.1	2.2	7.5
50% ben ^a	28	2.2	7.3
50% 3-pyr ^a	28	6.75	19.4
50% THF ^a	23.05	5.9	20.5
50% TDO ^a	20.5	1.8	8.1
Microplastics			
PA-6 ³²	30.3	5.5	15.4
PS ⁵¹	34.5	6.1	15.0
HDPE ⁵³	25.9	0	0.0
PETE ⁵³	39.3	4.2	9.7

^a Estimated as an average of literature values for pure compositions.

polarity values for the four plastics are shown in Table 1.³⁴ Beyond the polarity criterion, crystallinity was another property of general interest. In our atomistic models, HDPE and PETE are crystalline at standard temperature and pressure, PA-6 is amorphous, and PS is atactic and therefore amorphous. These microstructures are characteristic of MP waste.⁸

A range of adhesive compositions was created to bind MPs, with polyacrylates containing 2-EH side chains as the baseline case (Chart 1). This polymer was chosen because it was shown to bind MPs under aqueous conditions.⁵⁰ Derivatization by side-chain substitution should lead to different adhesive properties, which may enable selective MP binding. From Table 1 the percent polarities of acrylates go from 7.3% for poly(benzyl acrylate) (ben), which less than 2-EH at 7.5%, to 20.5% for THF. If polarity correlates with adhesive preference, polar adhesives should bind more strongly to polar MPs and *vice versa*. Both 3-pyridyl (3-pyr) and THF sidechains were chosen to form hydrogen bonds (H-bonds), an important consideration for aqueous adhesion. As plastics like PA-6 form H-bonds between amide groups, H-bonding acrylates could substitute for intra- and interchain H-bonds and enhance surface adhesion by more than would be expected from polarity considerations alone. Similarly, 3-pyr and ben have aromatic rings, which could boost adhesion to similar MPs (*e.g.*, PS) *via* pi-stacking. Finally, the C–F bonds of poly(tridecafluorooctyl acrylate) (TDO) is more hydrophobic,⁵⁶ which could enhance binding to the non-polar MPs (*e.g.*, HDPE) compared to polar ones. Together with the four distinct MPs, and the five side-chains under consideration, we anticipated differences in adhesion. Snapshots of the equilibrated MP-adhesive structures are in Fig. 1. Because the MP and adhesive atomistic models are of similar size, the surface contacts are mostly geometrically flat. Further analysis reveals that side-chain clustering should not be a concern in these surface contacts, as the side-chain composition sampling was performed randomly (SI, Section 2).

Aqueous work-of-adhesion

Aqueous WoA was determined through MD simulations for all four MP types combined with the acrylate copolymers. Fig. 1 shows the mean $WoA_{(aq)}$ for a given adhesive composition interacting with each MP, whereas Fig. 2 provides the WoA as



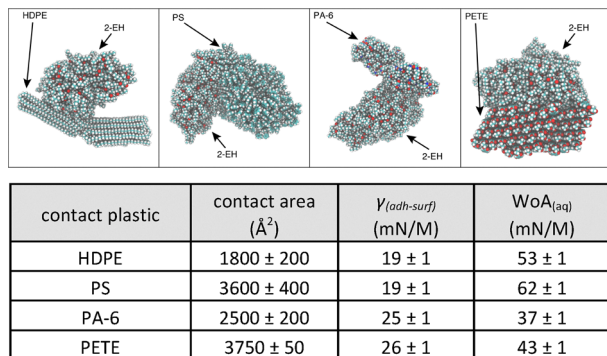


Fig. 1 (Top) 100% 2-EH adhesive interacting with microplastics. (Bottom) Contact area, average interfacial surface tension, and aqueous work of adhesion for the same surfaces. Averages were determined by periodic sampling of MD simulations with error from standard deviation.

a function of degree of substitution. For copolymers containing ben, THF, and 3-pyr side chains, the WoA increases with degree of substitution compared to 100% 2-EH (Fig. 2A, C and D), meaning that these side chains bind more strongly than the 100% 2-EH baseline.¹⁹ In contrast, the trend in WoA reverses with TDO substitution, showing a decreasing trend (Fig. 2B). Of the four side chains, THF resulted in the largest WoA_(aq) for the four MP types. While all four tested polyacrylates exhibit a preference for PS over other MP types, only TDO shows increasing selectivity with TDO composition. This preference can be seen in Fig. 2B, where at 50% TDO composition the WoA_(aq) gap between PS and HDPE has increased from ~ 10 mN m⁻¹ to ~ 20 mN m⁻¹.

Overall, the results can be used to check the hypotheses from the previous section. First, the degree of interaction between MP and adhesive is uncorrelated to adhesive side-chain aromaticity, as the slopes for PS and PETE are similar to HDPE and PA-6 for

the ben-substituted acrylate. Similarly, the hypothesized H-bonding between THF and 3-pyr with PA-6 was not observed, as evidenced by PA-6 remaining the lowest WoA_(aq) regardless of adhesive composition. Similarly, no correlation between percent polarity and WoA_(aq) was found (nor between the individual polar and dispersive surface energy values that determine percent polarity). For example, PS and PA-6 have similar % polarity values, yet the WoA_(aq) values are quite different regardless of the polyacrylate tested. Similarly, the more polar THF-based polymer exhibits the highest WoA_(aq) for all tested MPs irrespective of their polarity. Quantitative comparisons of % polarity *versus* WoA_(aq) can be found in the SI (Fig. S6–S9).

Structural changes of adhesives in aqueous conditions

To better understand how adhesives perform in water, several qualitative and quantitative metrics related to adhesion were examined. The equilibrated polyacrylate structures with 50% R* side chain compositions in the gas phase are compacted to maximize internal dispersive and polar interactions and minimize surface exposure (Fig. S10). Such behavior is generally expected under vacuum as there are no stabilizing surface interactions between vacuum and polymer. As seen in Fig. 3A, the polyacrylates undergo significant changes in the aqueous phase. For example, the TDO-substituted polyacrylate is still relatively compact whereas the THF-substituted polyacrylate is highly disrupted in aqueous simulations (*e.g.*, see hole formation). Similar deformations can be observed for 3-pyr, though to a lesser degree (Fig. S10). The TDO and ben-substituted polyacrylates distort in shape but still maintain somewhat compacted geometries (Fig. S6). These qualitative morphological changes suggest that there may be substantial differences in surface-area-to-volume ratios (SA:V) among different adhesives. Indeed, there was a quite significant increase in surface area for all adhesives relative to their initial conformations, as polymers containing 50% THF, TDO, 3-pyr and ben increased by $\sim 27\%$, $\sim 15\%$, 24% , and 16% , respectively, under aqueous conditions (Fig. S6).

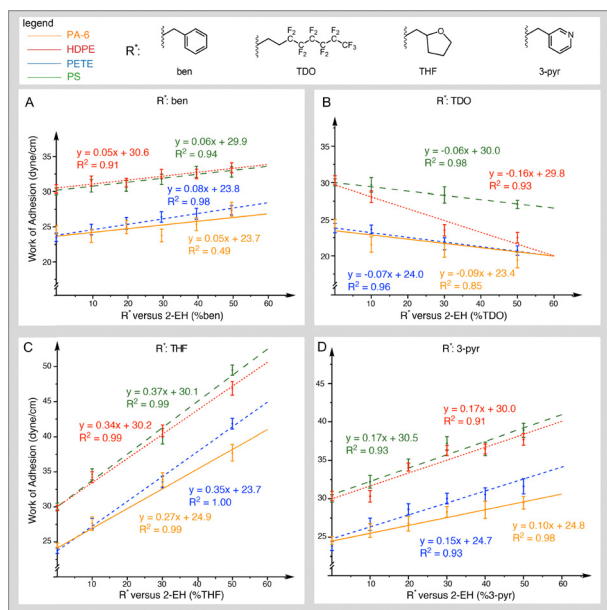


Fig. 2 WoA_(aq) results by R* percent composition compared to microplastic surfaces. (A) ben, (B) 3-pyr, (C) THF, (D) TDO.

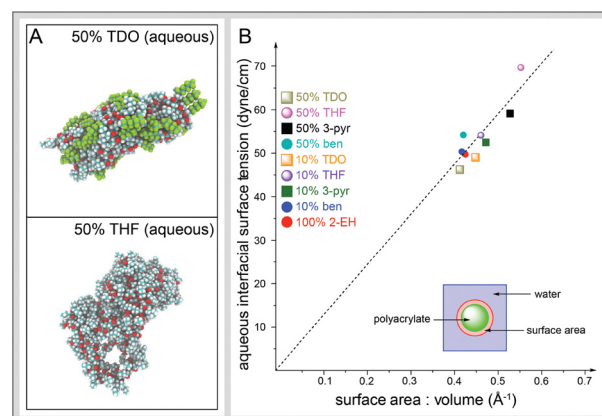


Fig. 3 (A) Representative structures of in aqueous simulation structures for 50% TDO (top) and THF (bottom) polyacrylates after simulation. (B) Interaction area *versus* $\gamma_{(adh-H_2O)}$ for representative polyacrylate adhesives, intercept is set to zero as non-interacting surfaces have no interaction energy.



Further examining the relationship between SA:V and $\gamma_{(\text{adh-H}_2\text{O})}$ is worthwhile to explain changes to the adhesives occurring in the aqueous phase. We found that the $\gamma_{(\text{adh-H}_2\text{O})}$ increases as SA:V increases, as highlighted in Fig. 3B. While all the materials follow roughly the same relationship between SA:V and $\gamma_{(\text{adh-H}_2\text{O})}$ ($R^2 = 0.997$), there are nuances worth noting. For instance, H-bond acceptor acrylates have significant changes in SA:V and $\gamma_{(\text{adh-H}_2\text{O})}$ compared to non-H-bond acceptor acrylates. The H-bond acceptor 3-pyr acrylate at 50% R^* composition shows 25% increases in SA:V and $\gamma_{(\text{adh-H}_2\text{O})}$ over the 100% 2-EH case. Conversely, for ben, which does not have an H-bond acceptor, the trend is somewhat steeper, as SA:V increases only $\sim 2\%$ at 50% R^* , with a 10% increase in $\gamma_{(\text{adh-H}_2\text{O})}$. This result indicates that H-bond accepting acrylates undergo greater morphological changes in the aqueous phase than non-acceptors. In contrast to the other three acrylate types, polymers with TDO sidechains decrease in SA:V and in $\gamma_{(\text{adh-H}_2\text{O})}$ with increasing % R^* . This trend can be attributed to the highly hydrophobic fluorocarbons, which resist contact with water. Overall, there is a notable SA:V correlation to $\gamma_{(\text{adh-H}_2\text{O})}$, and H-bonding appears to play a significant role in this relationship.

As hypothesized, H-bonds substantially influence polyacrylate behavior under aqueous conditions. Shown in Fig. 4A is the H-bond formation percentage (see SI, Section 10 for details on how H-bonds were assigned) for THF and 3-pyr side chains, which describes the ratio of H-bonds present compared to the total possible number of H-bonds for a given % R^* composition. This metric is not a measure of H-bonding strength, but rather the quantity. In the model, THF H-bonds were longer and weaker than their 3-pyr counterparts (Fig. 4A), which is expected given the weaker H-bond accepting strength of an ether. This finding is consistent with trends in Fig. 3B, since H-bonding between water and polyacrylate side chains is

generally more favorable than water interacting with the non-polar polymer backbone. This energetic driving force also leads to increasing disruption of the compact, non-aqueous morphology of polyacrylates as more H-bonds form, resulting in higher SA:V. The somewhat fewer number of H-bonds for 3-pyr relative to THF-substituted polymers is also consistent with this phenomenon, where the 3-pyr polyacrylates have lower $\gamma_{(\text{adh-H}_2\text{O})}$ than THF polymers of the same % R^* composition. Near non- (or weakly) polar surfaces, water exists in an ordered structure without strong interactions to the surface.⁵⁷ The water significantly rearranges when in contact with THF- or 3-pyr-containing polymers through formation of intermolecular H-bonds. The kernel density estimate (KDE) plots—similar to the pairwise distribution function but non-parametric to not bias the statistical analysis⁵⁸—shown in Fig. 4B quantify that water molecules form H-bonds at a median distance of 1.95 Å of THF and 3-pyr side chains and range in length from 1.5–2.5 Å.

There is also an increase in density of water molecules in the 2.5–3.0 Å range for the non-polar carbon and hydrogen atoms in THF and 3-pyr. In contrast, few water molecules are found for 2-EH, ben, and TDO in the same range. As the water molecules form H-bonds to the acrylate, interfacial surface tension increases, rather than decreases, as non-polar regions comprise the majority of polyacrylates. Ultimately, the driving force for increasing $\text{WoA}_{(\text{aq})}$ with H-bond accepting comonomer substitution is partly accounted for by disruption of the organized water shells, which increases interactions between water and non-polar regions. This disruption forces water molecules closer to the nonpolar regions of the same adhesive, which increases the surface tension and $\text{WoA}_{(\text{aq})}$. These relationships are contained in just one variable of the $\text{WoA}_{(\text{aq})}$ equation, so interactions between MPs and polyacrylate adhesives need to be examined next.

Properties of adhesive-plastic interactions

The determined $\gamma_{(\text{adh-plas})}$ are approximately one-half the magnitude of $\gamma_{(\text{adh-H}_2\text{O})}$, due to the similarity in surface polarity between MPs and polyacrylates, resulting in low interfacial tension (Fig. S13). For acrylates with ben, 3-pyr and THF sidechains, $\gamma_{(\text{adh-plas})}$ to the four MP types increases by 4–5 mN m^{-1} at 50% R^* composition compared to the 100% 2-EH case. These values compare to changes in $\gamma_{(\text{adh-H}_2\text{O})}$ of 10–20 mN m^{-1} for the same acrylates. For TDO-containing polymers, HDPE increases by $\sim 7 \text{ mN m}^{-1}$ across % R^* compared to the reference case (Fig. S13). At 50% TDO, $\gamma_{(\text{adh-plas})}$ for HDPE, PA-6, and PETE are all around $\sim 25 \text{ mN m}^{-1}$ while PS remains $\sim 19 \text{ mN m}^{-1}$ regardless of % R^* . These results show that $\gamma_{(\text{adh-H}_2\text{O})}$ is a strong factor for predicting trends in $\text{WoA}_{(\text{aq})}$ for TDO-containing polymers, while $\gamma_{(\text{adh-plas})}$ is more important for differentiating PS. This substantiates that the qualitatively different $\text{WoA}_{(\text{aq})}$ trend for PS originates from the interaction of PS and TDO where the $\gamma_{(\text{adh-plas})}$ remains low across % TDO, rather than from $\gamma_{(\text{adh-H}_2\text{O})}$ interactions.

In all, while $\gamma_{(\text{adh-H}_2\text{O})}$ is quantitatively more important to $\text{WoA}_{(\text{aq})}$, selectivity in MP adhesion is not tuned by adhesive interactions with water. Therefore $\gamma_{(\text{adh-plas})}$ remains a key consideration, despite its observed lower variance with adhesive composition.

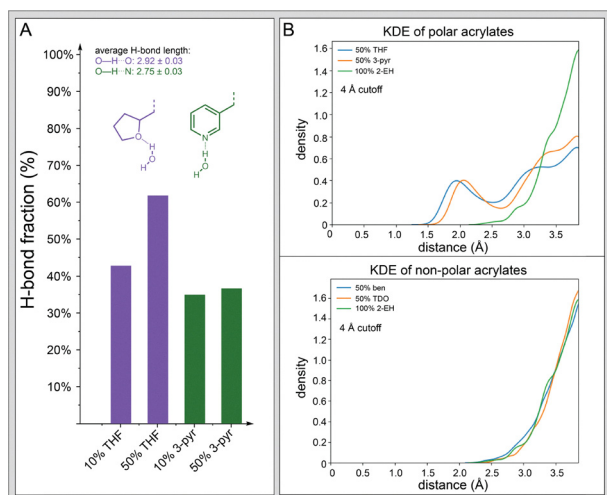


Fig. 4 (A) Normalized H-bond formation for representative polyacrylate adhesives (B). KDE of water molecules distance from sidechains (sampled within 4.0 Å of sidechains) for aqueous simulations of 50% polyacrylate composition compared against 100% 2-EH. Top is polar sidechains; bottom is non-polar sidechains.



Analysis of emergent behaviors in adhesion

The unique trends in $\gamma_{(\text{adh-plas})}$ for TDO warrant further examination to reveal their physical origin. Recall that while the contact area varies for each MP (Fig. 1), nearly all contacts are “flat” meaning that there is no obvious interweaving. This behavior is observed across almost all tested polyacrylate compositions. (Further images of flat surface contacts can be found in SI, Fig. S15–S26). The important exception to the observed flat contact occurs for TDO and PS (Fig. 5), where interweaving of 50% TDO is observed with PS but not HDPE. The degree of interweaving can be quantified by comparing the distance between the MP “centerline” and the “center” of TDO side chains. The KDE plots for 50% R^* repeat units interacting with PS show that most repeat units only approach ~ 10 Å of the centerline, whereas TDO has density near 3.5 Å from the PS centerline. This partial interweaving occurs because $\gamma_{(\text{adh-plas})}$ is lower than γ_{adh} and γ_{plas} , which creates a driving force for increasing interaction between them. The effect can be traced to increased surface tension of the acrylate due to TDO substitution.

Analysis of interweaving for TDO sidechain *versus* % composition (Fig. S14) indicates that increasing TDO from 10% to 50% increases the acrylate surface tension to the point of interweaving. This trend also provides evidence that TDO side chains are increasing the surface tension of the polyacrylate with increasing % TDO. Given that PA-6, HDPE, and PETE all have higher $\gamma_{(\text{adh-plas})}$ than PS, the surface tension of 50% TDO must be between

19–25 mN m^{-1} . To summarize, when $\gamma_{(\text{adh-plas})}$ is less than the surface tension of both the adhesive and MP, increased adhesion and interweaving can occur. These results show promise in achieving the goal of potentially separating PS from a mixture of the four tested plastics using TDO (or a similarly behaved) side chain.

Summary of properties that control adhesion ability

The above analysis revealed key properties that govern the overall behavior of polyacrylate adhesives. The H-bond accepting ability of the acrylate substituents was found to improve aqueous adhesion for THF and 3-pyr to all tested MPs. This effect was driven by the rearrangement of the ordered shells of water molecules surrounding non-polar surfaces. This disruption brings water molecules closer to non-polar regions in addition to forming H-bonds. As a result, the aqueous interfacial tension for H-bond accepting polyacrylates increases which, per eqn (1), increases $\text{WoA}_{(\text{aq})}$.

While ben, 3-pyr, and THF sidechains improved adhesive ability, the fluorocarbon TDO was found to reduce it. The overall decrease in adhesive strength for TDO polyacrylates is a result of their lower interfacial tension with water, where acrylate-water contacts are minimized. TDO polyacrylates also showed a change in binding selectivity to PS compared to the other MPs. This effect is caused by TDO's $\text{WoA}_{(\text{aq})}$ towards PS staying higher than for other MPs across changing side chain composition (% R^*). The unusual trend in $\text{WoA}_{(\text{aq})}$ for PS and HDPE originates from the interweaving between adhesive and MP; this was traced to the $\gamma_{(\text{adh-plas})}$ for 50% TDO binding to PS being less than γ_{adh} and γ_{plas} . This interweaving was a unique effect for TDO among the adhesive types tested in this work and may be a useful property for design of new selective acrylates.

Experimental comparison

An important consideration in developing novel adhesives is the difference between thermodynamic (from simulations) and practical adhesion (from experiments).⁵⁶ Practical adhesion is often orders of magnitude larger than thermodynamic adhesion.⁵⁶ The relationship between the two is typically assumed to follow:

$$G_C = \text{WoA}_{(\text{aq})} \cdot A(1 + \psi(a, T, \dots)) \quad (2)$$

Where G_C is the practical adhesive strength, γ_{adh} is thermodynamic adhesion (hereafter referred to as $\text{WoA}_{(\text{aq})}$), A is the area of interaction, and $\psi(a, T, \dots)$ is an empirical dissipation function, which depends on the adhesive and substrate composition, as well as experimental conditions.⁵⁶ In the case where $\psi(a, T, \dots) = 0$, G_C reduces to the thermodynamic adhesion. Interweaving or cross-linking contribute to $\psi(a, T, \dots)$ and tend to increase the magnitude of G_C .

Fig. 6 shows G_C from probe-tack measurements of 100% 2-EH and 42% ben with probes of the four MP compositions in aqueous conditions. Ben substitution was chosen as it would have a strong increase in $\text{WoA}_{(\text{aq})}$, relative to 100% 2-EH while having a smaller effect on $\psi(a, T, \dots)$, as they are more similar to 2-EH than the polar acrylates, THF and 3-pyr, as well as the fluorinated TDO. Overall, the probe-tack measure of practical adhesion replicates the general trends of the computed $\text{WoA}_{(\text{aq})}$. Uncertainties in the measurements remain, however, stemming

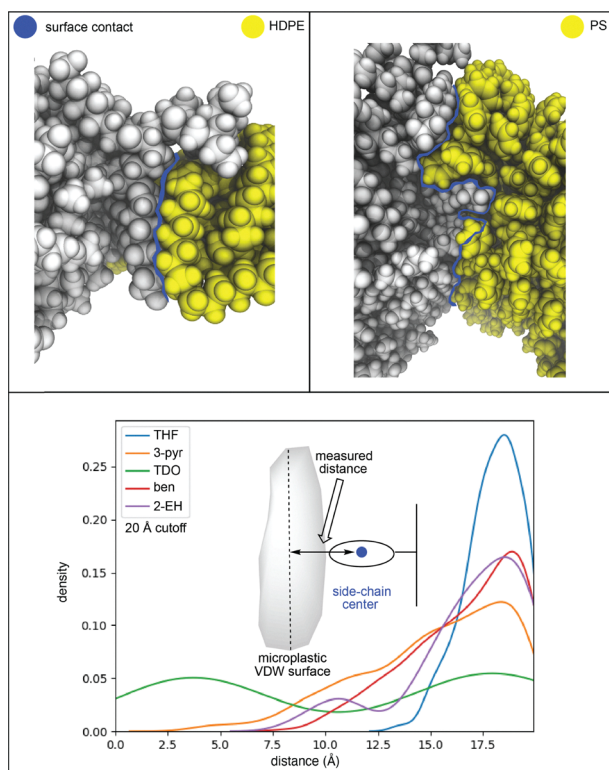


Fig. 5 (Top) Surface contacts between 50% TDO (white) and microplastics HDPE and PS (yellow). (Bottom) KDE plot of adhesive side chain distance (defined as the average cartesian coordinates for all atoms in a side chain) from PS centerline (defined as a line parallel to the contact between MP and adhesive that includes the volumetric center of the MP).



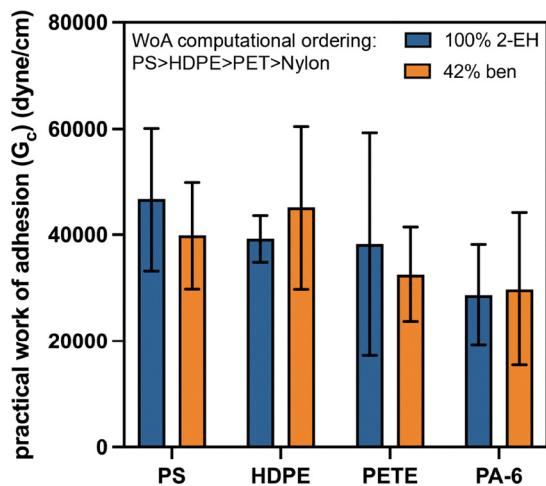


Fig. 6 Experimental Probe-tack comparison for select polyacrylate adhesives. Experiments were performed under aqueous conditions.

from the measurement being taken in water with plastic probes. For example, plastic probes will have differences in surface roughness that affect the practical $WoA_{(aq)}$ through changes in probe contact area. Keeping these uncertainties in mind, comparisons can still be made between G_c and computed WoA. The lowest G_c values come with PA-6, which matches the computational results. PS, PETE, and HDPE have similar values of G_c and are also the highest in computational $WoA_{(aq)}$. While comparisons between ben-substituted adhesives with 100% 2-EH adhesives could further test the computed trends, these differences could not be resolved within the error bars of the experiments.

Nevertheless, this work demonstrates that computationally determined values of thermodynamic adhesion are substantially more predictive of practical adhesion than comparing percent polar compositions (see SI, Fig. S6–S9), which represents a significant advance in available tools for developing adhesives.

Conclusions

While H-bonding produced a strong shift in aqueous interaction energy, the hypothesized H-bonding between nylon and H-bond-accepting acrylates (3-pyr, THF) was not observed. Along the same lines, aromatic acrylates, with ben and 3-pyr side chains, did not show π -stacking with the aromatic polystyrene. Instead, the interfacial energies between the four MPs and adhesives with ben, THF, and 3-pyr sidechains remained relatively consistent in both value and trend (Fig. 2). This consistency limited the viability of using these three adhesives for selective capture of MPs. Adding perfluoro sidechains, however, was found to increase selectivity for PS. This behavior arose from an increase in aqueous/side-chain surface energies with increasing substitution, which eventually surpassed the interfacial energy between TDO and PS. When this occurs, the interface becomes more favorable than wetting both surfaces, driving interweaving between TDO and PS. The result was an emergent phase not seen in the other adhesive-MP interactions. This phenomenon may warrant further attention in future studies.

The concepts formulated in this work may be useful to precisely tailor the next generation of polyacrylate adhesives for MP binding. In particular, the microscopic features, such as H-bond accepting in aqueous adhesion, intra- and inter-polymeric interactions, and interweaving, should all be considered when selecting an adhesive polymer design. Highly selective adhesives might be achieved by further reducing interfacial energies to a specific MP and increasing overall surface energy of polyacrylates. The computational tools presented here are well-suited to further refining mechanistic understanding in search of this behavior. Besides these concepts, atomistic modeling can likely be used to predict WoA for novel acrylate compositions, further expanding the scope of knowledge in this area. The inability to fully model all components of practical adhesion remains a limitation, preventing a complete determination of experimental adhesive utility. In all, this understanding and these tools can be used to design improved polyacrylate adhesives.

Author contributions

T. M. E. J.: methodology, formal analysis, writing – original draft. H. E. T.: formal analysis, investigation, writing – review & editing. M. T. R.: conceptualization. W. J.: methodology, investigation. M. E. C.: investigation. A. J. M.: resources, writing – review & editing, supervision, project administration, Funding acquisition. P. M. Z.: conceptualization, resources, writing – review & editing, supervision, project administration, funding acquisition.

Conflicts of interest

There are no conflicts to declare.

Data availability

Additional data is available from the corresponding author upon reasonable request.

Supplementary information (SI): further computational details, polyacrylate monomer sequences, experimental methods, tables, and figures can be found in the supplemental information. See DOI: <https://doi.org/10.1039/d5cp03631c>.

Acknowledgements

The authors acknowledge NSF EFRI E3P Grant 2029251 through the National Science Foundation–Emerging Frontiers in Research and Innovation 2020 program for support of this work.

References

- 1 E. Penzel, *Ullmann's Encyclopedia of Industrial Chemistry*, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, Germany, 2000.
- 2 T. Ohara, T. Sato, N. Shimizu, G. Prescher, H. Schwind, O. Weiberg, K. Marten, H. Greim, T. D. Shaffer and P. Nandi, *Ullmann's Encyclopedia of Industrial Chemistry*, Wiley, 2020.
- 3 N. Xu, J. Cao and X. Liu, *J. Macromol. Sci., Part B: Phys.*, 2015, **54**, 1153–1168.



- 4 Y. Peykova, O. v Lebedeva, A. Diethert, P. Müller-Buschbaum and N. Willenbacher, *Int. J. Adhes. Adhes.*, 2012, **34**, 107–116.
- 5 P. T. Chazovachii, J. M. Rieland, V. V. Sheffey, T. M. E. Jugovic, P. M. Zimmerman, O. Eniola-Adefeso, B. J. Love and A. J. McNeil, *ACS ES&T Eng.*, 2021, **1**, 1698–1704.
- 6 D. H. Kaelble, *J. Appl. Polym. Sci.*, 1974, **18**, 1869–1889.
- 7 J. P. G. L. Frias and R. Nash, *Mar. Pollut. Bull.*, 2019, **138**, 145–147.
- 8 J. Barrett, Z. Chase, J. Zhang, M. M. B. Holl, K. Willis, A. Williams, B. D. Hardesty and C. Wilcox, *Front. Mater. Sci.*, 2020, **7**, 576170.
- 9 M. Padervand, E. Lichtfouse, D. Robert and C. Wang, *Environ. Chem. Lett.*, 2020, **18**, 807–828.
- 10 J. Boucher and D. Friot, Primary microplastics in the oceans: A global evaluation of sources, IUCN International Union for Conservation of Nature, 2017.
- 11 M. Bergmann, S. Mützel, S. Primpke, M. B. Tekman, J. Trachsel and G. Gerdt, *Sci. Adv.*, 2019, **5**, 1157.
- 12 M. R. Michielssen, E. R. Michielssen, J. Ni and M. B. Duhaime, *Environ. Sci.*, 2016, **2**, 1064–1073.
- 13 M. Wang, P. Zhou, S. DuBay, S. Zhang, Z. Yang, Y. Wang, J. Zhang, Y. Cao, Z. Hu, X. He, S. Wang, M. Li, C. Fan, B. Zou, C. Zhou and Y. Wu, *J. Hazard. Mater.*, 2025, **487**, 137274.
- 14 Y. Xiang, Z. Wang, Y. Zhao, J. Liu, J. Wang, Q. Lu and L. Xie, *J. Colloid Interface Sci.*, 2025, **683**, 347–356.
- 15 A. Chamas, H. Moon, J. Zheng, Y. Qiu, T. Tabassum, J. H. Jang, M. Abu-Omar, S. L. Scott and S. Suh, *ACS Sustain. Chem. Eng.*, 2020, **8**, 3494–3511.
- 16 E. Schmaltz, E. C. Melvin, Z. Diana, E. F. Gunady, D. Rittschof, J. A. Somarelli, J. Virdin and M. M. Dunphy-Daly, *Environ. Int.*, 2020, **144**, 106067.
- 17 G. Casoli and S. Ramkumar, *Mare Plasticum – The Plastic Sea*, Springer International Publishing, Cham, 2020.
- 18 J. Hopewell, R. Dvorak and E. Kosior, *Philos. Trans. R. Soc., B*, 2009, **364**, 2115–2126.
- 19 J. R. Dann, *J. Colloid Interface Sci.*, 1970, **32**, 302–320.
- 20 C. A. Finch, Polymer Handbook, 3rd edn, *Br. Polym. J.*, 1990, **23**, 277–280.
- 21 W. J. Shim, S. H. Hong and S. Eo, *Microplastic Contamination in Aquatic Environments*, Elsevier, 2018.
- 22 A. Kowalski, Z. Czech and L. Byczyński, *J. Coat. Technol. Res.*, 2013, **10**, 879–885.
- 23 L. W. McKeen, *Film Properties of Plastics and Elastomers*, Elsevier, 2012.
- 24 A. J. Kinloch, *Adhesion and Adhesives*, Springer, Netherlands, Dordrecht, 1987.
- 25 D. Briggs, D. G. Rance and B. J. Briscoe, *Comprehensive Polymer Science and Supplements*, Elsevier, 1989.
- 26 J. J. Bikerman, *Surface Chemistry: Theory and Applications*, Elsevier, Burlington, 2nd edn, 2013.
- 27 M. K. Abdelrahman, H. Kim, J. Maeng, P. Ondrusek and T. H. Ware, *Macromolecules*, 2020, **53**, 2388–2395.
- 28 M. A. Firestone, S. C. Hayden and D. L. Huber, *MRS Bull.*, 2015, **40**, 760–767.
- 29 C. W. Abney, S. Das, R. T. Mayes, L.-J. Kuo, J. Wood, G. Gill, M. Piechowicz, Z. Lin, W. Lin and S. Dai, *Phys. Chem. Chem. Phys.*, 2016, **18**, 23462–23468.
- 30 T. Khudiyev, E. Huseyinoglu and M. Bayindir, *Sci. Rep.*, 2014, **4**, 4607.
- 31 R. G. Winkler, J. Elgeti and G. Gompper, *J. Phys. Soc. Japan*, 2017, **86**, 101014.
- 32 L. Xie, F. Liu, J. Liu and H. Zeng, *ACS Appl. Mater. Interfaces*, 2020, **12**, 58360–58368.
- 33 L. Xie, X. Cui, J. Liu, Q. Lu, J. Huang, X. Mao, D. Yang, J. Tan, H. Zhang and H. Zeng, *ACS Appl. Mater. Interfaces*, 2021, **13**, 6941–6950.
- 34 S. Ebnesajjad, *Surface Treatment of Materials for Adhesive Bonding*, Elsevier, 2014.
- 35 K. O'Driscoll and R. A. Sanayei, *Macromolecules*, 1991, **24**, 4479–4480.
- 36 J. A. McCammon, B. R. Gelin and M. Karplus, *Nature*, 1977, **267**, 585–590.
- 37 M. Levitt and S. Lifson, *J. Mol. Biol.*, 1969, **46**, 269–279.
- 38 S. A. Hollingsworth and R. O. Dror, *Neuron*, 2018, **99**, 1129–1143.
- 39 N. V. Bhagavan, *Medical Biochemistry*, Elsevier, 2002.
- 40 B. Park, M. Chandross, M. J. Stevens and G. S. Grest, *Langmuir*, 2003, **19**, 9239–9245.
- 41 K. F. Mansfield and D. N. Theodorou, *Macromolecules*, 1991, **24**, 4295–4309.
- 42 D. L. Cheung, D. P. McMahon and A. Troisi, *J. Phys. Chem. B*, 2009, **113**, 9393–9401.
- 43 D. M. Huang, R. Faller, K. Do and A. J. Moulé, *J. Chem. Theory Comput.*, 2010, **6**, 526–537.
- 44 K. N. Schwarz, T. W. Kee and D. M. Huang, *Nanoscale*, 2013, **5**, 2017–2027.
- 45 D. Alberga, G. F. Mangiatordi, L. Torsi and G. Lattanzi, *J. Phys. Chem. C*, 2014, **118**, 8641–8655.
- 46 M. Moreno, M. Casalegno, G. Raos, S. V. Meille and R. Po, *J. Phys. Chem. B*, 2010, **114**, 1591–1602.
- 47 C. Poelking and D. Andrienko, *Macromolecules*, 2013, **46**, 8941–8956.
- 48 O. Alexiadis and V. G. Mavrantzas, *Macromolecules*, 2013, **46**, 2450–2467.
- 49 S. L. Mayo, B. D. Olafson and W. A. Goddard, *J. Phys. Chem.*, 1990, **94**, 8897–8909.
- 50 S. Jo, T. Kim, V. G. Iyer and W. Im, *J. Comput. Chem.*, 2008, **29**, 1859–1865.
- 51 Y. Li, J. Q. Pham, K. P. Johnston and P. F. Green, *Langmuir*, 2007, **23**, 9785–9793.
- 52 S. N. Omenyi, A. W. Neumann and C. J. van Oss, *J. Appl. Phys.*, 1981, **52**, 789–795.
- 53 D. K. Owens and R. C. Wendt, *J. Appl. Polym. Sci.*, 1969, **13**, 1741–1747.
- 54 M. S. Hanqi Wen, Johns Hopkins University, 2019.
- 55 *CRC Handbook of Chemistry and Physics*, ed. W. M. Haynes, CRC Press, 2014.
- 56 P. Kirsch, *Modern Fluoroorganic Chemistry*, Wiley-VCH, Weinheim, 1st edn, 2004, vol. 1.
- 57 D. A. Dillard, *Advances in Structural Adhesive Bonding*, Elsevier, 2010, pp. 350–388.
- 58 M. Kooi and A. A. Koelmans, *Environ. Sci. Technol. Lett.*, 2019, **6**, 551–557.

