

# Photo-Induced Shrinking of Aqueous Glycine Aerosol Droplets

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**ABSTRACT:** Due to their small size, micrometer and submicron sized solution droplets can respond differently to physical and chemical processes compared with extended bulk material. Using optically trapped micrometer sized aqueous glycine droplets, we demonstrate a photo-induced degradation of glycine upon irradiation with visible light, even though molecular glycine does not absorb light in the near UV/vis range to any significant extent. This reaction is observed as photo-induced shrinking of the droplet, which we characterize by analyzing the elastic light scattering and the Raman spectrum of the droplet over the course of the reaction. We find the volume to shrink with a constant rate over the major part of the shrinking process. This indicates the presence of a rate limiting photocatalyst, which we attribute to mesoscopic glycine clusters in the droplet solution. Our findings relate to previous reports about enhanced absorption and fluorescence rates of amino acid solutions. However, to the best of our knowledge, this is the first experimental evidence of a photochemical pathway facilitated by mesoscopic clusters. Light interaction with such mesoscopic photoactive molecular aggregates might be more important for aerosol photochemistry than previously anticipated.

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## 1 1. INTRODUCTION

2 Aerosols, dispersions of solid and liquid particles in a gas, are ubiquitous in Earth's atmosphere and as such play  
3 an important role for many atmospheric processes (Boucher et al., 2013; PöSchl and Shiraiwa, 2015). Size, chem-  
4 ical composition, viscosity and thermodynamic phase of aerosol particles respond to their environment including  
5 the surrounding gas species, temperature, humidity (Bones et al., 2012; Zieger et al., 2017; Tang et al., 1997;  
6 Swietlicki et al., 2008) and light irradiation (PöSchl and Shiraiwa, 2015; Cremer et al., 2016; Walser et al., 2007;  
7 Corral Arroyo et al., 2022). Particular attention has been paid to their chemistry distinct from that in the bulk, a  
8 phenomenon possibly arising from high surface to volume ratio of aerosol particles and the accessibility to highly  
9 supersaturated states (Altaf et al., 2016; Kucinski et al., 2019; Bzdek and Reid, 2017). Various chemical reactions  
10 have been shown to be accelerated in microdroplets (Cremer et al., 2016; Lee et al., 2015; Girod et al., 2011),  
11 with some reactions being exclusive to the droplet phase (Lee et al., 2019). These unique reactors can act as  
12 medium for the birth, growth and degradation of atmospherically relevant particles (Ruiz-Lopez et al., 2020), and  
13 be utilized for organic synthesis (Bain et al., 2017). Chemical processes in prebiotic aerosols have also been  
14 proposed as potential mechanisms for the origin of life (Tervahattu et al., 2004). However, molecular processes  
15 leading to the anomalous chemistry in micrometer and submicron aerosol particles are still largely unknown.  
16 Although some processes may be ascribed to the discontinuous and asymmetric intermolecular interactions at the  
17 particle surface (Ruiz-Lopez et al., 2020), the microphysical origins behind many of the aforementioned particle  
18 specific phenomena are still not adequately explored.

19  
20 Glycine is an amino acid that acts as precursor to proteins and fulfills a number of other biological functions  
21 (Arnstein, 1954; Hall, 1998; Jackson, 1991). With its small size and simple structure, glycine often serves as a  
22 proxy for other amino acids and physiologically relevant molecules, and as such has been studied extensively in  
23 the past. It is generally accepted that molecular glycine does not absorb light in the near UV/vis range, similar to  
24 other amino acids (Bhat and Dharmaparakash, 2002). However, it has been shown that their optical properties  
25 change when glycine molecules arrange themselves into mesoscopic clusters **by forming hydrogen bond networks**  
26 (Terpugov et al., 2021). Furthermore, these formations respond to light irradiation in non-trivial ways which can  
27 be exploited to induce long range order inside the glycine solution (Alexander and Camp, 2019; Sugiyama et al.,  
28 2012; Zaccaro et al., 2001; Garetz et al., 2002; Urquidi et al., 2022) on a scale of up to millimeters (Yuyama et  
29 al., 2010). While these interactions have the potential to change the optical properties of glycine ensembles sig-  
30 nificantly and to enable new photochemical reaction pathways, there has been little experimental evidence for  
31 such reactions so far.

32  
33 In this work, we study the response of aqueous glycine droplets to irradiation by visible light. We observe the  
34 shrinking of optically trapped micrometer sized glycine droplets, which can be unambiguously attributed to the  
35 exposure to the trapping laser with wavelength 532 nm. To the best of our knowledge, this interaction has not  
36 been reported before. We characterize it here with particular focus on the shrinking rate and its dependence on the

37 light intensity. To explain our results, we discuss possible reaction schemes based on the available experimental  
38 data. Although further data is needed to elucidate the exact photochemical pathways of the observed reaction,  
39 these findings demonstrate the existence of a photochemical reaction for molecules which previously have been  
40 considered photochemically inert at visible wavelengths.

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42

## 43 2. METHODS

44 Dual beam optical traps are widely used to confine and isolate single particles (Ashkin, 1997; Gong et al., 2018;  
45 Čižmár et al., 2005; Esat et al., 2018; Reich et al., 2020). The counter-propagating tweezers (CPT) setup for  
46 trapping aqueous glycine droplets is shown in Fig. 1 and consists of a continuous green laser beam (Novanta  
47 Photonics Opus 532 6W), which is expanded and then split into two beams of equal power. These two beams are  
48 aligned counter-propagating on a single axis and focused into the trapping cell, where a single droplet is trapped  
49 between the two focii.

50

51 The droplets are generated from 1.0 M or 2.0 M aqueous solutions of glycine (purity 100%, HPLC certificate,  
52 Sigma-Aldrich G7126) using a commercial atomizer (TSI 3076) with pressurized, humidified nitrogen gas (purity  
53 5.0). We performed additional elementary and HPLC analyses of the glycine and the glycine solutions. The results  
54 showed no indication of any trace impurities absorbing in the visible range. On this basis, we conclude that our  
55 samples do not contain any contaminants that could act as photosensitizer for photochemical reactions (see be-  
56 low). A system of copper tubings directs the spray of particles into the trapping cell, where the droplets agglom-  
57 erate at the designated trapping position. The humidification of the nitrogen flow is necessary to ensure that the  
58 droplets reach the trapping position in the liquid state. The trapping cell is filled with nitrogen gas (purity 5.0) and  
59 a steady nitrogen flow formed by combining wet and dry nitrogen with adjustable flow ratios is used to control  
60 the relative humidity (RH) in the cell. For the experiments reported here, the RH is set at  $77 \pm 3$  % well above the  
61 efflorescence RH of glycine at approximately 55 % (Chan et al., 2005). At this RH, the droplet solution is super-  
62 saturated with an estimated glycine concentration of 60 % in mass (Chan et al., 2005), corresponding to approxi-  
63 mately 5 M. Temperature and RH inside the trapping cell are monitored by a sensor (Sensirion SHT35) placed a  
64 few millimeters away from the trapping position. When the agglomerated particle reaches a size of approximately  
65 2-3 $\mu$ m in radius, the remainder of the droplets in the cell are flushed out with nitrogen for 20-30 minutes to ensure  
66 that only the trapped droplet remains in the cell. After flushing, the power of the trapping laser is kept constant  
67 until the end of the measurement.

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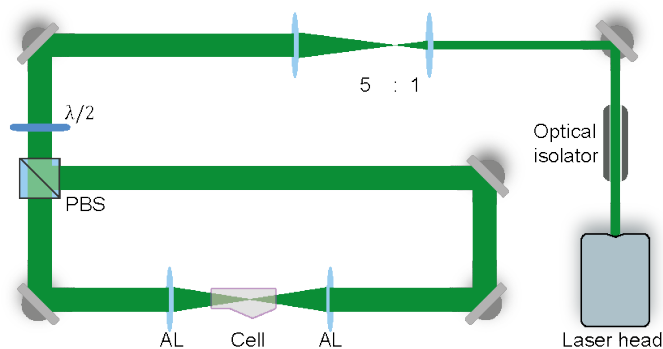
69 The particle shrinking is monitored by imaging the polarization resolved two-dimensional angular optical scatter-  
70 ing (polarization resolved TAOS) of the particle (Parmentier et al., 2022), as shown in Fig. 2. The TAOS image  
71 is obtained by collecting the elastically scattered light of the trapping beams under a scattering angle of  $90 \pm 24^\circ$   
72 with an objective (Mitutoyo 20x NA 0.42). The parallel and perpendicular polarization components with respect  
73 to the scattering plane (TAOS PPol and TAOS SPol) are separated using a polarization beam splitter and recorded  
74 with separate CMOS cameras (Thorlabs DCC1545M). The scattering intensity for each polarization is calculated  
75 from the average of the respective TAOS image and recorded over time. At specific times, the shrinking spherical  
76 particle reaches a size at which it is in resonance with the light of the trapping beams, which corresponds to a Mie  
77 resonance (Bohren and Huffman, 2008). From the comparison of the recorded evolution of the polarization re-  
78 solved scattering intensity to simulations using Mie theory, the size of the particle can be determined at the specific

79 times. Fig. 3 shows an example of such a TAOS analysis. The values of the size at the discrete points in time,  
80 obtained from the times where the particles experience a Mie resonance, can then be interpolated with high accu-  
81 racy to obtain the full size evolution of the particle over the course of the measurement.

82

83 The molecular composition of the particle is monitored by continuous recording of Raman spectra (David et al.,  
84 2020) during the shrinking process. To this end, the light scattered by the particle is collected under a scattering  
85 angle of  $90 \pm 24^\circ$  by a second objective and fiber coupled into a low noise, high sensitivity spectrograph (Andor  
86 KY-328i-A). The inelastically scattered light is analyzed in the range 540-680 nm which corresponds to Raman  
87 shifts of 280-4100  $\text{cm}^{-1}$ . This range contains in particular the O-H symmetric stretching mode of water ( $\nu_2\text{-H}_2\text{O}$ ,  
88 2700-3750  $\text{cm}^{-1}$ ) as well as several vibrational modes of glycine, which we exploit for the characterization of the  
89 molecular composition (see later data for an example).

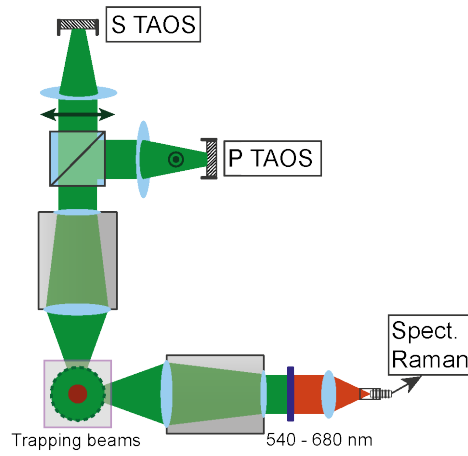
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92 **Figure 1** Counter-propagating tweezers setup. The laser beam is first expanded by a factor of 5 and then split into  
93 two beams by the polarizing beam splitter (PBS). The half-waveplate ( $\lambda/2$ ) rotates the polarization to  $45^\circ$  with  
94 respect to the axes of the PBS to ensure equal power splitting between the two beams. The beams are aligned on  
95 a single axis and focused into the trapping cell using two aspherical lenses (AL). An optical isolator introduced  
96 at the start of the beam path prevents unwanted optical feedback into the laser head.

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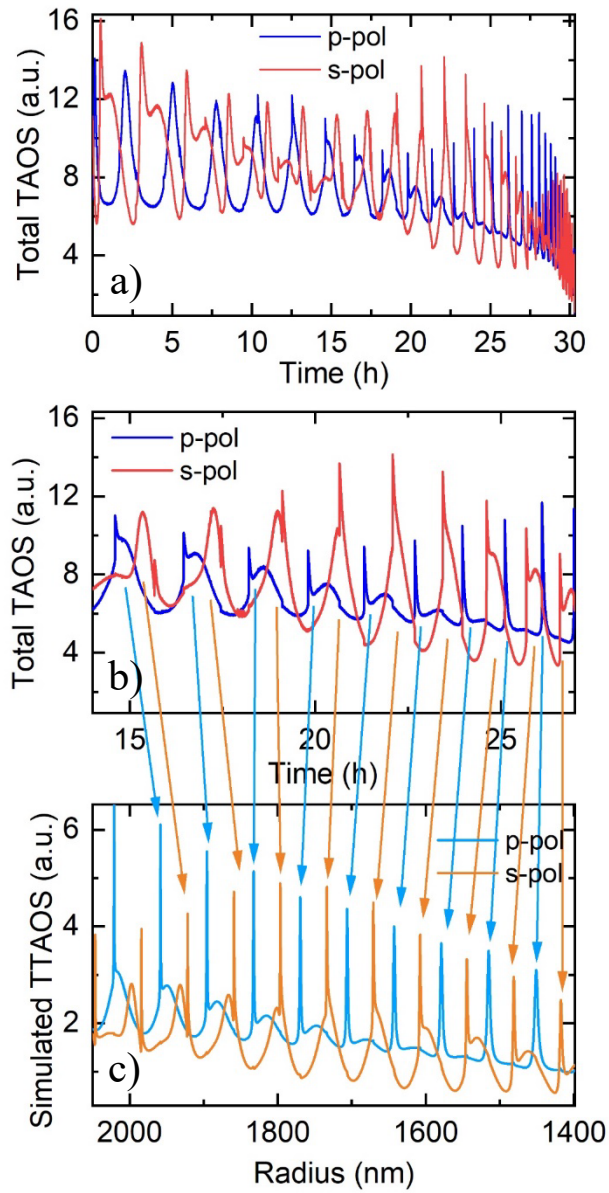
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**Figure 2** Setup for TAOS imaging and Raman spectroscopy. The trapping beams run perpendicular to the figure plane. The scattered light of the trapping beams is collected horizontally and vertically at a scattering angle of  $90 \pm 24^\circ$ . The vertical beam is split into parallel and perpendicular polarized light with respect to the scattering plane and the respective beam is loosely focused on a CMOS camera (P TAOS and S TAOS respectively). The horizontal beam is filtered for the spectral range of 540 – 680 nm and fiber coupled into a low noise, high sensitivity spectrometer (Spect. Raman) for measurement of the Raman spectrum.



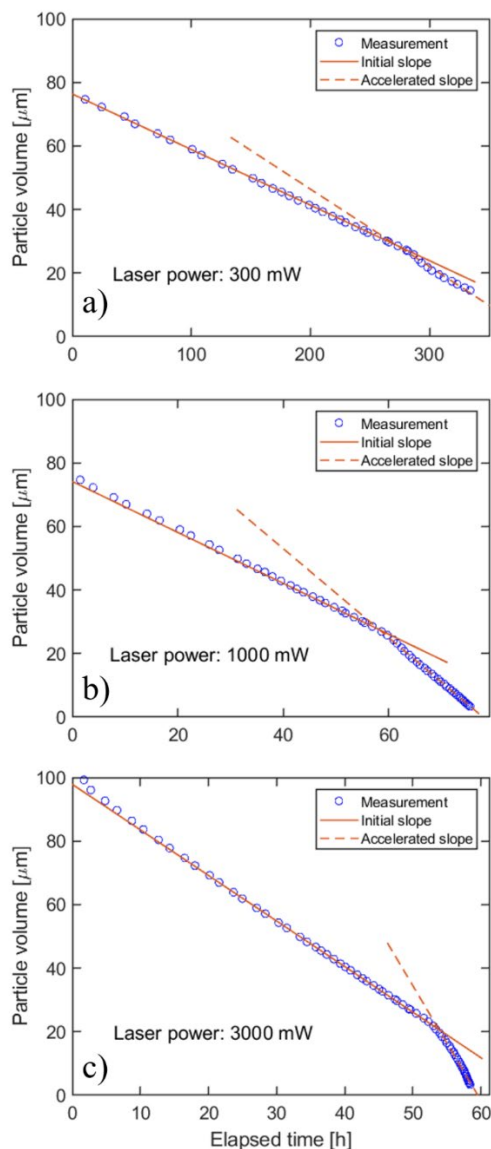
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107 **Figure 3** Analysis of the two-dimensional angular optical scattering spectrum (TAOS). **a)** Experimental polari-  
 108 zation resolved total TAOS signal over time. **b)** Zoom on a specific time interval for clarity **c)** Simulated polari-  
 109 zation resolved total TAOS signal as function of particle radius in the specific time interval. Arrows indicate the  
 110 peak assignment based on the similarities between the peak shapes.

111

112 **3. RESULTS AND DISCUSSION**

113 The shrinking of the aqueous glycine droplets over time is shown in Fig. 4 for three representative examples. The  
114 droplets are trapped using different laser powers, which affects the rate at which their volume is decreasing. For  
115 all laser powers, the volume is observed to shrink linearly with time over a large portion of the shrinking process.  
116 After a certain point (‘point of acceleration’), when the droplet has lost 60 – 80 % of its volume, the shrinking  
117 suddenly accelerates, only to continue again approximately linearly with time, albeit at a higher rate.

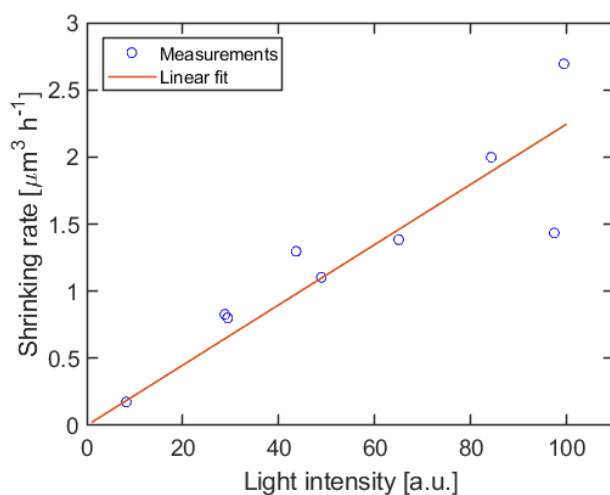


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119 **Figure 4** Volume shrinking of glycine droplets as a function of time. **a)** Droplet trapped at 300 mW nominal laser  
120 power **b)** droplet trapped at 1000 mW nominal laser power **c)** droplet trapped at 3000 mW nominal laser power.  
121 At higher power, the shrinking proceeds faster (visible by the larger slopes). Solid and dashed lines indicate the  
122 best linear fit before and after the acceleration of the shrinking observed at approximately **a)** 280 h, **b)** 59 h and  
123 **c)** 53 h.



124 To quantify the dependence of the shrinking rate on the light intensity incident on the particle, a linear fit is  
125 performed on the data before and after the point of acceleration. Since between different experiments, the align-  
126 ment of the optical trap, and hence the focusing of the laser light on the particle, is subject to temporal mechanical  
127 drifts, the nominal laser power used for trapping of the droplets is not an optimal indicator for the incident inten-  
128 sity. Instead, we use the intensity of the scattered light as a measure that is proportional to the incident light  
129 intensity. The intensity of the scattered light is obtained from the average signal of the TAOS PPol and SPol  
130 images during the time of the droplet shrinking. To ensure consistency between the different experiments, the  
131 average of the light intensity is taken over the same volume interval of  $[65.4, 51.0] \mu\text{m}^3$  (radius  $[2.5, 2.1] \mu\text{m}$ ) for  
132 all droplets. This interval corresponds to the interval for which we obtained data for most droplets. Fig. 5 shows  
133 the resulting initial shrinking rates as function of light intensity. The shrinking rate is observed to be proportional  
134 to the light intensity, confirming that the shrinking is induced by the trapping laser at a wavelength of 532 nm.



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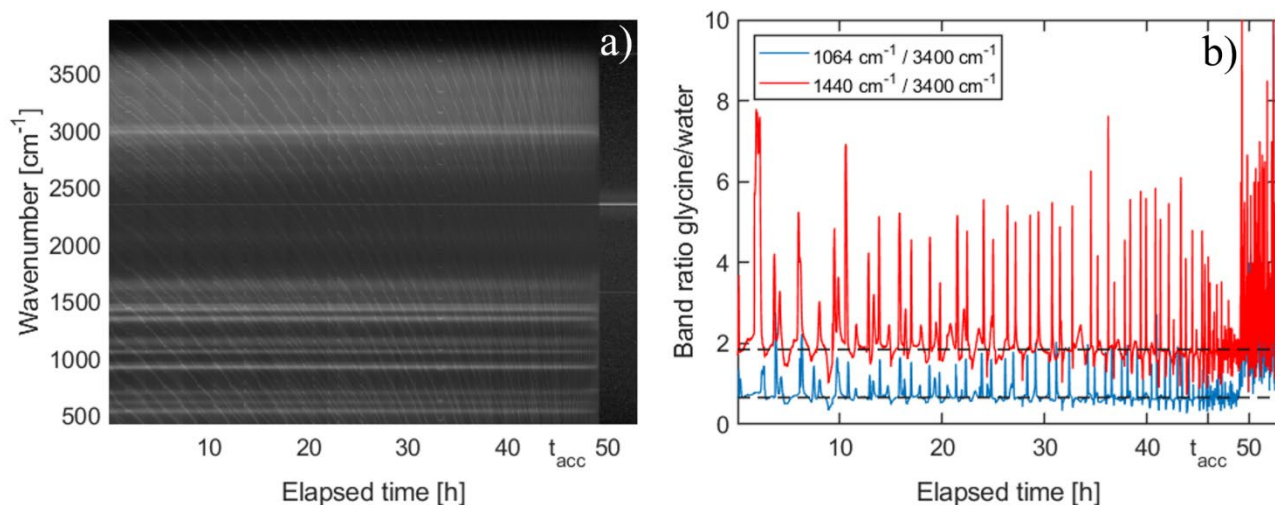
136 **Figure 5** Fitted shrinking rate as function of light intensity. The solid line indicates the best linear fit through  
137 the origin.

138 To gain further insight into the shrinking mechanism, we analyze the evolution of the Raman spectra over the  
139 course of the shrinking process. A representative example of such an evolution is shown in Fig. 6. For Raman  
140 spectra of single spherical particles, so called morphology dependent resonances, or whispering gallery modes  
141 (WGMs) (Oraevsky, 2002), are superimposed on the molecular signals. To separate the WGMs from the molecular  
142 signal of interest, the Raman spectra are normalized and stacked in chronological order from left to right, as shown  
143 in Fig. 6a. The WGMs show up as a manifold of thin slanted lines bending towards lower wavenumbers with  
144 increasing time as the droplet shrinks. Molecular band positions on the other hand are independent of particle  
145 size, and are identified as horizontal lines in the evolution of the Raman spectra.

146

147 From Fig. 6a it is evident that the molecular Raman signal remains qualitatively the same, indicating that no  
148 significant change in the molecular composition takes place in the droplet over the course of the shrinking process.  
149 This behavior is observed for all investigated droplets. Since the particle loses the major part of its volume during

150 the shrinking, this implies that glycine is removed from the droplet as a consequence of chemical reaction (*vide*  
 151 *infra*). As the water vapor pressure of the droplet is given by the surrounding RH of 77 % and therefore has to  
 152 remain constant, the removal of glycine from the droplet must be accompanied by the evaporation of water in  
 153 order to maintain the equilibrium glycine concentration. A quantitative analysis of the Raman spectrum (Fig. 6b)  
 154 reveals that the spectral intensity of glycine modes with respect to the O-H stretching mode of water remains  
 155 constant, confirming a constant concentration of glycine molecules during the shrinking process. The measure-  
 156 ment ends when the particle becomes too small for stable trapping, and therefore leaves the optical trap.  
 157



158 **Figure 6** Temporal evolution of Raman signal during droplet shrinking. **a)** Normalized Raman spectra stacked  
 159 chronologically from left to right. The molecular Raman bands are visible as horizontal straight white lines on the  
 160 dark background. The finer, slanted and curved lines correspond to whispering gallery modes. After approxi-  
 161 mately 49 h, the particle leaves the optical trap as it reaches a size that is too small for trapping, and only back-  
 162 ground is recorded. The band remaining afterwards at 2323 cm<sup>-1</sup> corresponds to the nitrogen gas in the trapping  
 163 cell. **b)** Ratio of the molecular glycine bands centered at 1064 cm<sup>-1</sup> and 1440 cm<sup>-1</sup> to the water band at around  
 164 3400 cm<sup>-1</sup>. Peaks and dips in the graphs correspond to spectra where a whispering gallery mode is superimposed  
 165 on the glycine signal and the water signal, respectively, and are not relevant for the molecular composition. The  
 166 dashed horizontal lines are a guide to the eye. In this example, the acceleration of the droplet shrinking is observed  
 167 at  $t_{acc} = 46$  h. From this point in time onwards until the particle is lost, the faster shrinking leads to more frequent  
 168 WGMs perceived as an apparent increase of noise in the data.  
 169

170 Aqueous droplets which contain a stable non-volatile solute do not shrink over time when they are optically  
 171 trapped (see Supplementary Section S1 for an example). The observation of droplet shrinking in the case of dis-  
 172 solved glycine proves that glycine does not remain stable inside the droplet solution. Furthermore, the linear  
 173 dependence of the shrinking rate on the light intensity (Fig. 5) proves that the observed shrinking is induced by  
 174 the laser light. In particular, the volume shrinking rate increases from 0.18 to 2.25  $\mu\text{m}^3 \text{h}^{-1}$  when the light intensity

175 is increased by the same relative amount, thus spanning over one order of magnitude. This observation is intri-  
176 guing as aqueous glycine is not known to absorb light in the visible range, similar to other amino acids (Bhat and  
177 Dharmaparakash, 2002). In addition, the linear dependence between shrinking rate and light intensity rules out the  
178 possibility of multiphoton absorption, which might otherwise populate energetically excited states of glycine to  
179 induce chemical reactions. The fact that the observed shrinking is driven by the incident light indicates a photo-  
180 chemical reaction or photothermal processes occurring in the aqueous glycine droplet. As is shown in the follow-  
181 ing however, photothermal processes can be ruled out by our experimental data.

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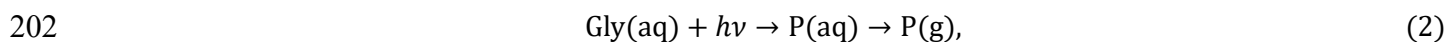
183 The observation of a constant shrinking rate over a large portion of the experiments (Fig. 4) provides another  
184 important piece of information, as this allows us to rule out some of the possible shrinking mechanisms. For  
185 instance, assuming some residual absorption of light at 532 nm by the droplet solution, one might argue that the  
186 shrinking is a consequence of enhanced evaporation due to the droplet heating by the laser. The evaporation of  
187 glycine from the droplet can be approximated by the Hertz-Knudsen equation:

$$188 \quad \frac{dN}{dt} = S \cdot \frac{\alpha p}{\sqrt{2\pi MRT}} \quad (1)$$

189 where  $\frac{dN}{dt}$  is the molar evaporation rate of glycine,  $S$  is the droplet's surface,  $p$  is the partial pressure and  $M$  is the  
190 molar mass of glycine,  $R$  the gas constant,  $T$  the temperature and  $\alpha$  a heuristic sticking coefficient with values  
191 between 0 and 1. It is evident from Eq. (1) that the rate of shrinking by evaporation should scale with the surface  
192 area of the droplet, and that in this case a deceleration of the shrinking should be observed over time, contrary to  
193 the experimental data. Moreover, any heating by laser light absorption would at most lead to a very small temper-  
194 ature rise due to the efficient cooling by the surrounding gas (see Supplementary Section S2 for an upper estimate  
195 of the temperature increase). In addition to these arguments, if evaporation were significant, some evaporation  
196 should still be observable even at low light intensity, where the heating of the droplet is negligible and hence any  
197 deviations from room temperature can be neglected. As seen from Fig. 5 however, there is no shrinking observable  
198 for low light intensities. We can therefore rule out evaporation as the dominant shrinking mechanism.

199

200 The observation of a constant shrinking rate also excludes photochemical reactions in which molecular glycine  
201 directly absorbs photons:

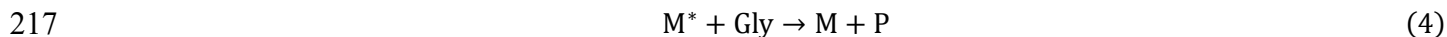
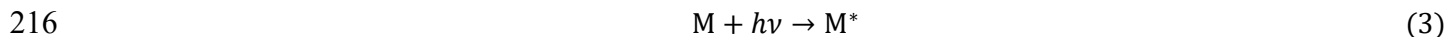


203 where Gly(aq) is a solvated glycine molecule,  $h\nu$  is the energy of the incoming photon and P is the reaction prod-  
204 uct that is removed from the droplet (aq) into the surrounding gas phase (g) afterwards. As mentioned above,  
205 molecular glycine is considered non-absorbing at 532 nm. However, if we nevertheless assume that molecular  
206 glycine could be very weakly absorbing as in Eq. (2), the photon density inside the droplet would be constant over  
207 the course of the photochemical reaction. Earlier studies (Parmentier et al., 2021; Corral Arroyo et al., 2022) show  
208 that no significant size dependence of the optical confinement is observed for weakly absorbing particles in the  
209 micrometer- and submicrometer-size ranges. Hence, the first step in Eq. (2) would be pseudo-first order, for which

210 the reaction rate is proportional to the concentration of glycine molecules in the droplet. Therefore, one would  
211 expect a decrease in the observed shrinking rate over time, which contradicts the experimental observation.

212

213 The examples above illustrate that any mechanism, in which glycine molecules directly absorb incoming photons,  
214 cannot explain the constant shrinking rates. This implies that a more intricate reaction must take place in the  
215 droplet. We suggest the following simplified scheme with an additional reaction partner M:



218 where  $M^*$  denotes a photoexcited state of M, and P is the reaction product of glycine in the presence of this pho-  
219 toexcited species. This scheme represents a mediated reaction in which the reaction partner M is activated by light  
220 absorption, and then reacts with glycine and returns back to the ground state. M shows the characteristics of a  
221 photosensitizer (Corral Arroyo et al., 2018; George et al., 2015; Wang et al., 2020; Rapf and Vaida, 2016), which  
222 is not consumed during the reaction, and therefore the amount of M remains constant in the droplet. We further  
223 assume that the light absorption of the photosensitizer M is the rate limiting step, i.e., that Eq. (3) proceeds much  
224 slower than Eq. (4). This is equivalent to requiring that M is only weakly absorbing, or that the concentration of  
225 M in the droplet is low. Since the amount of M in the rate limiting step (Eq. (3)) remains constant during the  
226 shrinking process, so does  $M^*$  (quasistationary), resulting in a constant rate of degradation of Gly (Eq. (4)). As-  
227 suming that the absolute concentration of M is much smaller than that of Gly at all times, and therefore has no  
228 relevant effect on the equilibrium water vapor pressure of the droplet, the volume shrinking rate is predicted to be  
229 constant in accordance with the experimental data.

230

231 Although the proposed mechanism in Eq. (3) and (4) concurs with the observed constant shrinking rates, it does  
232 not yet specify the nature of the photosensitizer and its reaction with glycine. We first discuss potential candidates  
233 for the photosensitizer. Contamination during the preparation of the different aqueous glycine solutions used in  
234 this study was minimized by using pure substances (glycine purity  $\geq 99\%$ , water resistivity  $18.2 \text{ M}\Omega\cdot\text{s}$ ). No  
235 correlation between the shrinking rates in Fig. 5 and the age of the solution at the time of the measurements was  
236 observed, which indicates that there is no accumulation of photoactive contaminants in the solution after the  
237 preparation. We therefore argue that contamination is not the origin of the photosensitizer.

238

239 It is evident from the previous discussion that the photosensitizer must possess an absorption band at 532 nm,  
240 which is not the case for single solvated glycine molecules. The optical properties of molecules may change  
241 however when forming intermolecular bonds, e.g. leading to an enhancement of the absorption and fluorescence  
242 in the case of protein aggregates (Homchaudhuri and Swaminathan, 2004; Shukla et al., 2004; Chan et al., 2013;  
243 Pinotsi et al., 2013) and amino acid clusters (Chen et al., 2018). For glycine solutions in particular, observations  
244 of light absorption in the near UV/vis range have been attributed to the presence of mesoscopic clusters (Jawor-

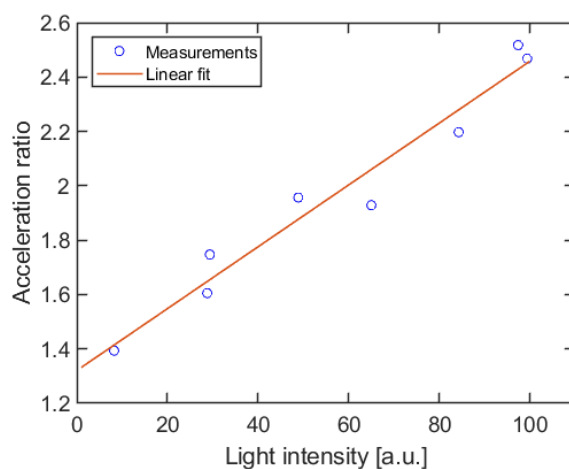
245 Baczynska et al., 2013; Zimbitas et al., 2019), specifically due to the formation of hydrogen bonds between indi-  
246 vidual molecules (Terpugov et al., 2021).

247  
248 Mesoscopic clusters occur naturally in both undersaturated and supersaturated glycine solutions, though a kinetic  
249 barrier may have to be overcome for their formation (Jawor-Baczynska et al., 2013). ). A similar barrier needs to  
250 be overcome to dissolve these clusters. Hence the clusters can remain kinetically stable even if the number of  
251 glycine monomers decreases. The average size of the mesoscopic clusters, typically of the order of 100 nm for  
252 bulk solutions, depends not only on the monomer concentration but also on the history of the sample, indicating  
253 that the clusters do not necessarily remain in thermodynamic equilibrium after formation (Jawor-Baczynska et  
254 al., 2013). Based on these observations, we propose that the photo-induced droplet shrinking is mediated by light  
255 absorption of mesoclusters in the glycine solution. These mesoclusters remain kinetically stable during the droplet  
256 shrinking, despite the decrease of the absolute number of glycine monomers (i. e. constant glycine monomer  
257 concentration). Hence, these clusters act as stable photosensitizers inside the droplets, in accordance with the  
258 observed constant shrinking rate. The observation of a constant initial shrinking rate thus allows us to narrow  
259 down the possible reaction mechanisms at work in the droplets. Alternative schemes might be conceivable, such  
260 as the existence of several reaction partners for glycine. However, there is no further experimental evidence in  
261 favor of more complex alternatives to the simple mechanism proposed (Eq. (3) and (4)). Furthermore, as pointed  
262 out above, mesoscopic glycine clusters match the requisite characteristics of the proposed photosensitizer, and are  
263 therefore likely candidates. To pursue the argument, let us further discuss the role of the mesoscopic clusters in  
264 the observed acceleration of the droplet shrinking.

265  
266 The acceleration of the droplet shrinking proceeds relatively promptly at the point of acceleration when the parti-  
267 cle has lost a typical amount of 75 % of its volume (Fig. 4). Assuming that the total number of mesoclusters  
268 remains approximately constant ( $M$  in Eqs.(3) and (4)), their concentration has increased by an approximate factor  
269 of 4 at this point. The sudden nature of the shrinking acceleration hints at a phase transition inside the particle. In  
270 particular, the increase in nanocluster concentration may trigger the separation of a dense cluster phase inside the  
271 droplets as part of a liquid-liquid phase transition. While a definite conclusion has to await more detailed micro-  
272 scopic investigations, we present the following arguments in favor of this explanation. Mesoscopic clusters in  
273 aqueous solutions are known to interact with focused light irradiation by assembling in the focal point of the light  
274 beam due to the optical force that acts on the individual clusters (Sugiyama et al., 2012). This mechanism is the  
275 basis of laser-induced phase transitions(Alexander and Camp, 2019), in the particular case of glycine both for  
276 liquid-liquid phase separation (Sugiyama et al., 2012; Yuyama et al., 2010) and solid crystal nucleation (Sugiyama  
277 et al., 2012; Alexander and Camp, 2019; Yuyama et al., 2010; Zaccaro et al., 2001; Garetz et al., 2002). It should  
278 be noted at this point that the spot size in the focus of our optical trap is slightly larger (5  $\mu\text{m}$ ) than the typical  
279 droplet size, and that therefore, it might appear unlikely that the electromagnetic field gradient is strong enough  
280 to induce cluster aggregation in our case. However, micrometer sized droplets exhibit a large variance in the  
281 spatial distribution of the internal light field, owing to the nanofocusing effect (Cremer et al., 2016; Corral Arroyo

282 et al., 2022), which can provide the field gradients necessary for aggregation. If in the case of our trapped droplets,  
283 the acceleration is due to a phase transition, it would likely be assisted by the light irradiance. One would therefore  
284 expect a dependence of the observed shrinking acceleration on the incident light intensity. Fig. 7 shows the meas-  
285 ured acceleration ratio, that is, the ratio between the shrinking rate after to before the point of acceleration, as a  
286 function of light intensity. From this data it is evident that the ratio increases with higher light intensity, which  
287 agrees with our explanation of a light-induced phase transition. Since in this scheme, the clusters are expected to  
288 aggregate in the regions of high light intensity after reaching a critical concentration, this would lead to a larger  
289 absorption rate and thus a larger subsequent reaction rate, in agreement with the observation. Further studies will  
290 be necessary to provide conclusive evidence for this explanation.

291



292

293 **Figure 7** Acceleration of droplet shrinking versus light intensity. The solid line represents the best linear fit  
294 through the measurement data.

295 More data will also be required to understand the specifics of the interaction between the photosensitizer and the  
296 solvated glycine molecules in Eq. (4). Here, we can only provide a qualitative discussion based on the available  
297 data. The Raman spectra (Fig. 6) show no detectable change in the molecular composition, even after the droplet  
298 has lost the major part of its volume during the shrinking process. Since this observation rules out the accumula-  
299 tion of reaction products in the droplets over time, the reaction products must be small, volatile compounds that  
300 quickly evaporate into the surrounding gas phase. Known degradation mechanisms of glycine and other amino  
301 acids in aqueous solution proceed via reaction with radical species, in particular solvated electrons  $e_{\text{solv}}^-$  and hy-  
302 droxyl  $\cdot\text{OH}$  radicals (Moenig et al., 1985; Garrison, 1964, 1972), which form as part of a photosensitized reaction  
303 with chromophoric organic matter in water (Lundeen et al., 2014; Sun et al., 2018; Mopper and Zika, 1987).  
304 Currently, our data does not allow us to distinguish between different degradation pathways.

305

#### 306 4. CONCLUSIONS

307 We have demonstrated that single micrometer sized aqueous glycine droplets respond to the illumination with  
308 laser light of 532 nm by shrinking, despite the fact that molecular glycine does not absorb in the near UV/vis

309 range. Most remarkably, the volume shrinking rate remained constant over the major part of the shrinking process.  
310 This indicates a photo-induced decay of glycine molecules in the presence of a rate limiting catalyst, or photosen-  
311 sitizer, and the subsequent evaporation of small, volatile reaction products. Based on the available literature data,  
312 we propose that intrinsic mesoscopic clusters of glycine molecules formed by hydrogen bonding in the aqueous  
313 solution are the most plausible candidates for this photosensitizer. The presence of mesoscopic glycine clusters  
314 would also explain the sudden acceleration of the shrinking rate occurring at a volume loss of ~75 %. Because of  
315 its dependence on the light intensity, we attribute this sudden rate change to arise from the interaction of the  
316 mesoclusters with the incident light, possibly initiating a light-induced phase transition.

317

318 This study provides yet another example of the non-trivial interactions of light with aqueous glycine solutions  
319 (Alexander and Camp, 2019; Sugiyama et al., 2012; Zaccaro et al., 2001; Garetz et al., 2002; Yuyama et al., 2010),  
320 which facilitate previously undiscovered reaction pathways - interactions that are likely not exclusive to glycine.  
321 Light harvesting by and light interaction with such mesoscopic photosensitizers in aerosol droplets might also  
322 have played a role in the formation of more complex organic molecules under prebiotic conditions. Further in-  
323 vestigations are needed to shed light on the specifics of the observed phenomena, and to yield new insight into  
324 the underlying reaction mechanisms, which remain elusive in part. Studying solutions in micrometer sized drop-  
325 lets (attoliter volumes) using high laser powers offers the advantage of much higher sensitivity to photo-induced  
326 reactions than typically achievable with bulk solutions.

327

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335

## 336 **Data Repository**

337 The data that support our findings are deposited on the ETH Research Collection under <https://doi.org/xx>.

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