Dual-responsive nanoparticles that aggregate under the simultaneous action of light and CO$_2$†

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Metallic nanoparticles co-functionalised with monolayers of UV- and CO$_2$-sensitive ligands were prepared and shown to respond to these two types of stimuli reversibly and in an orthogonal fashion. The composition of the coating could be tailored to yield nanoparticles capable of aggregating exclusively when both UV and CO$_2$ were applied at the same time, analogously to the behaviour of an AND logic gate.

Stimuli-responsive nanomaterials† that self-assemble in a reversible fashion are important because of their emerging applications in diverse fields, such as switchable catalysis,2 water purification,3 detection of analytes,4 and time-sensitive information storage.5 Accordingly, a variety of nanoparticles (NPs) have been engineered to aggregate in response to different external stimuli, such as a magnetic field,7 pH,8,9 or chemical fuel.10 For example, dynamic aggregation of metallic NPs was accomplished by reversibly photo-switching the electronic states and geometries of surface-bound chromophores.11–14 New applications could arise, however, for NPs switching the electronic states and geometries of surface-bound amines, NHCs (N-heterocyclic carbenes), or guanidines,15–18 whereas they exhibited no detectable response to CO$_2$ (considering the recent report of a CO$_2$-sensitive organic polymer and functionalisation of metallic NPs with CO$_2$-responsive materials). Here, we presume that control of the NP surface polarity by using fine-tuned amounts of photo- and CO$_2$-sensitive ligands could induce selective and reversible aggregation of NPs with two orthogonal stimuli (Fig. 1).

It is well established that mixtures of organically functionalised long-chain alkanethiols can form mixed self-assembled monolayers (mSAMs) on the surfaces of metallic NPs (here, we focused on gold NPs). We commenced our study with AuNPs functionalised with mSAMs comprising an azobenzene-terminated thiol (AzoSH) and a “background” ligand, 1-dodecanethiol (C$_{12}$SH) (Fig. 2a). As the UV-Vis spectra in Fig. 2c show, NPs decorated with critical (50%) amounts of AzoSH aggregated in toluene upon exposure to long-wave (i = 365 nm) UV light,31 whereas they exhibited no detectable response to CO$_2$ (confirmed by dynamic light scattering (DLS) measurements; see ESI† Fig. S2 in Section S5). Furthermore, we observed that the trans-to-cis photoisomerisation of AzoSH did not affect the affinity of these NPs towards CO$_2$ (Fig. S2, ESI†). Therefore, we concluded that those AuNPs...

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† Electronic supplementary information (ESI) available: Synthesis and characterisation of ligand 3; synthesis and functionalisation of gold nanoparticles; and control experiments. See DOI: 10.1039/c4cc08541h

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**Fig. 1** (A) Reversible isomerisation of a photoswitchable ligand. (B) Reversible CO$_2$ complexation by a diamine ligand. (C) A dual-stimuli-responsive AND logic gate.
bearing only the AzoSH and C12SH ligands are inert towards the CO2 stimulus under these conditions.

The lack of general knowledge on the construction of CO2-sensitive metal-nanoparticle surfaces encouraged us to investigate the synthesis and application of thiol-terminated amine ligands (Fig. 1b). Based on previous literature reports, we presumed that ethylenediamine derivatives are capable of CO2 complexation under low-pressure conditions. To verify the effect of CO2 binding on the solubility in toluene, we synthesised and studied the behaviour of a model compound, N-decylethylenediamine. As detailed in the ESI† (Section S3), this diamine was readily soluble in toluene; however, rapid precipitation commenced upon bubbling CO2 through the solution (Fig. S1, ESI†). The process was reversible: the addition of an inert gas (e.g. N2) caused the precipitate to redissolve. Encouraged by these results, we then proceeded to synthesise AuNPs decorated with the diamine moieties. Although diamine-functionalised NPs were previously prepared by reacting 11-bromoundecane-1-thiol (2)-decorated NPs with 1,2-ethylenediamine, we began our investigation by preparing a well-defined thiol bearing the diamine moiety. As shown in Scheme 1, the synthesis of diamine 3 was straightforward, starting with a known material 1. With the diamine-functionalised thiolate ligand 3 in hand, we prepared a family of dual stimuli-responsive AuNPs by varying the ratio of the two responsive ligands (AzoSH and diamine 3) and the background ligand 1-dodecanethiol (C12SH), as shown in Fig. 3. Although 3 could potentially bind to AuNPs via both the mercapto and the amine functionalities, the latter scenario is unlikely given the ca. two-orders-of-magnitude higher affinity of Au to thiols as compared to amines.40 In addition, binding to Au via the amine functionalities would cause NP crosslinking and irreversible aggregation, which we did not observe.40

Owing to the high polarity and low toluene solubility of the diamine ligand 3, the ligand exchange reaction was conducted under dilute (ca. 10 times) conditions, compared with typical ligand exchange reaction conditions.41 Moreover, 3 was always used after drying under vacuum at 40 °C to ensure that it was not complexed with CO2. The functionalised AuNPs were precipitated with methanol (1:1 v/v) in the presence of didodecyldimethylamonium bromide (0.5 mg per 0.716 mg of Au), washed extensively with methanol to remove any unbound molecules, dried under vacuum and kept in an inert atmosphere prior to use. A good indication of the presence of 3 in the functionalised AuNPs is their good solubility in polar solvents (e.g., ethanol, methanol and even water), which confirms a successful ligand–exchange reaction (for comparison, AuNP–AzoSH is soluble only in non-polar solvents such as toluene, chloroform, or dichloromethane). The functionalisation procedure did not affect the size or size distribution of the NPs, whose monodispersity was confirmed by transmission electron microscopy (Fig. 5a).

Fine-tuning of the ratio of the three ligands was critical for obtaining NPs exhibiting the behaviour shown in Fig. 1c (in this context, it is worth mentioning that the previously adapted on-nanoparticle reaction between ethylenediamine and 2-functionalised AuNPs would hardly be applicable). We found that when the molar percentage of 3 in the NPs exceeded 25%, the particles were insoluble in toluene due to their high
surface polarity. These aggregated NPs could not be redissolved using visible light, sonication, or by purging with N₂ (to remove any possibly complexed CO₂). Therefore, the amount of diamine 3 was fixed at 25% throughout the optimisation of the mSAM composition. Having screened the ratio of the remaining two ligands (AzoSH and C₁₂SH) from 1:2 to 2:1, we found that NPs containing 33–35 mol% of AzoSH within the mSAM exhibited the behaviour shown in Fig. 1c. According to DLS, these NPs had hydrodynamic diameters of 7–10 nm in toluene, and did not aggregate upon either only UV irradiation or CO₂ treatment (bubbling into the solution) (purple and blue bars, respectively, in Fig. 4). Only when both stimuli were applied at once (4 min of UV irradiation during CO₂ bubbling) did the NPs rapidly assemble into aggregates of 800–1000 nm in size, as shown by DLS (green bars in Fig. 4). The presence of aggregated NPs was also evident from TEM and spectroscopic analysis (Fig. 5b and c). Interestingly, these aggregates were stable in a closed vial placed in the dark for more than 30 minutes.

Further demonstration of the AND gate-like behaviour was provided by removing one of the two stimuli. For example, when a solution of aggregated NPs was sonicated, facile disaggregation was observed even under constant UV irradiation (red bars in Fig. 4; ca. 10 nm). The resulting NPs did not respond to only CO₂ or UV, but they could again be quickly aggregated by applying UV and CO₂ simultaneously (second-cycle green bars in Fig. 4 and the TEM image in Fig. 5b). Interestingly, our NPs also assembled as a result of CO₂ bubbling (4 min), followed by exposure to UV light, or via the opposite sequence, whereby UV-irradiated NPs were treated with CO₂ in the dark: these results indicate that both (i) cis-azobenzene and (ii) the CO₂-diamine complex are relatively stable, (i) in the dark, and (ii) unless other stimulants (such as sonication, heating or inert gas bubbling) are applied.

Thus assembled AuNPs readily deaggregated under ambient light (5–6 minutes) or upon exposure to visible light (fluorescent bulb, <30 seconds), and consequently, additional assembly–disassembly cycles could be performed. Overall, we concluded that this study represents the first example of CO₂- and light-responsive nanoparticles, whose behaviour constitutes an AND logic gate. Noteworthy, NPs having higher surface concentrations of AzoSH did respond to UV light by assembling into aggregates, which could aggregate further upon the delivery of CO₂ (ESI,† Fig. S6C). In other words, only upon pre-exposure to UV light were these specific NPs responsive to CO₂.

We prepared gold nanoparticles co-functionalised with azobenzene- and diamine-terminated ligands, which rendered the particles responsive to UV irradiation and CO₂, respectively. Exposure to UV and CO₂ led to a stepwise increase in the nanoparticle surface polarity, triggering aggregation in a non-polar medium (toluene). The composition of the mixed monolayer of ligands could be tuned in order to afford NPs capable of aggregating only when both types of stimuli are applied at once. Both of these stimuli could easily be turned off/removed without the generation of any by-products/chemical waste, after which the assembly–disassembly cycles could be repeated. The aggregates exist only when both light and CO₂ are applied simultaneously. An important future direction concerning these and other man-made dynamically aggregating materials is to study/engineer their emergent behaviours under non-equilibrium conditions.

Notes and references
Under our experimental conditions, transformation of the carbamate into a cyclic urea is highly unlikely as it typically requires high CO pressures, elevated temperatures, and/or the presence of a catalyst (e.g. ref. 38 and 39).

This conclusion is supported by an experiment in which we incubated AuNPs co-functionalized with 3 and C12SH under oxygen for an extended period of time. Under oxidative conditions, the presence of free SH groups on the outer surfaces of NPs would likely cause crosslinking due to disulfide bridge formation. No such crosslinking was observed by DLS or UV-Vis spectroscopy.

In addition to the increasing NP surface polarity in the nonpolar medium, attractive forces between the NPs could originate from direct interactions (e.g. due to dipole–dipole interactions of the \( \text{cis-azobenzene dipoles} \)).