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Environmental and health risks of nanorobots: an early review

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Nanorobots for biomedical applications have experienced extensive research and rapid development during the last decade, up to a point where they can now deliver cargos to designated sites in organisms under laboratory conditions. Despite this development, research into nanorobot risks and discussions about potential regulation of nanorobots have so far been limited. This early review of risks related to nanorobots first provides a brief overview of the current state of the technology. The overview outlines three main types of nanorobots: helices, nanorods and DNA nanorobots. Several different designs exist for each of these categories. Second, early indications of potential hazards are reviewed and discussed. Two potential hazards are highlighted: (i) the use of hazardous materials and UV light in nanorobots, and (ii) the loss of propulsion/targeting control. Third, how current regulations are adapted to nanorobots is discussed. Current regulations for medical devices are clearly not adapted to nanorobots and it is even unclear which specific regulations might be applicable. In order to make the most of the use of nanorobots, we recommend they should be subject to broad, risk-related studies as well as dialogues with stakeholders and the public about the definition, purpose and controllability of nanorobot applications. A list of ten priority questions to be addressed in future risk-related studies of nanorobots is provided.

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Environmental significance

So far, discussions and research about environmental and health risks related to nanomaterials have focused on so-called passive nanomaterials, such as metal-oxide nanoparticles and carbon nanotubes. Much less attention has been given to so-called active nanomaterials. This includes nanorobots, which are individual nano-sized devices able to perform designated tasks, so far given limited attention regarding their potential future risks. This paper provides a review of early indications of potential environmental and health hazards related to nanorobots, investigates the applicability of existing regulatory frameworks, and provides recommendations on engaging expert stakeholders and the public.

1. Introduction

In a foundational paper, Roco¹ distinguished between a number of generations of nanotechnology, the two earliest being *passive* and *active* nanostructures. Tour² used a similar categorization and defined passive nanotechnology as when “the nano part does nothing particularly elaborate”. Active nanotechnology was defined as when “the nano entity does something elaborate such as absorbing a photon and releasing an electron, thereby driving a device, or moving in a specific and definable fashion across a surface”. Although the exact distinction between the passive and active nanotechnology can be tricky, conventional nanoparticles and nanotubes currently used in existing nanoproducts^{3,4} generally belong to the

category of passive nanomaterials. This is where most efforts in terms of risk-related research have occurred during the 2000s, in particular for a limited set of nanomaterials, including silver nanoparticles, titanium dioxide nanoparticles, silica nanoparticles, cerium dioxide nanoparticles, zinc oxide nanoparticles, iron nanoparticles, quantum dots, fullerenes, carbon nanotubes and graphene.^{5–9} Much less attention has been given to active nanomaterials, probably because of their limited production and use in society. However, one type of active nanomaterial is clearly on the march. Often referred to as science fiction, nanorobots are currently being extensively researched and developed, especially for medical applications where there is an effort to merge nanotechnology with pharmaceuticals.^{10,11} The most frequent application mentioned is drug delivery, in particular for site-specific cancer treatment through the delivery of tumor-killing drugs.¹² Other envisioned areas of applications beside medicine include environmental monitoring and water remediation.¹³

Despite this development, research on risks related to nanorobots have so far been limited. History shows several

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examples of how the introduction of new technology that offered great benefits into society later turned out to also cause notable environmental and health impacts.^{14,15} Regulation was generally imposed only at a late stage, long after the first signs of negative side effects appeared. A telling example from medicine is ionizing radiation, discovered around 1900, which brought important diagnostic and therapeutic benefits through X-rays and radioisotopes. Already at that time, there were early reports that exposure to radiation could cause severe damage, but these were largely overlooked. X-rays were even used for fitting shoes and removing unwanted hair in beauty shops in the 1930s and 1940s. Such misuses were possible because of the lack of regulation ensuring the safety of ionizing radiation. Only very slowly have such regulations evolved as the knowledge about radiation risks grew bigger, eventually enabling the beneficial use of radiation while keeping risks reasonably low.¹⁶ Another important medical discovery – perhaps the most important of all times – is antimicrobials, which have saved millions of lives by killing malignant bacteria. In parallel with such predominantly beneficial uses, antimicrobials have also been used as growth promoter in industrialized animal husbandry. Animals fed with sub-therapeutic amounts of antimicrobials for long time periods showed enhanced properties, such as increased growth rate, food conversion, egg production and milk yield. Much a consequence of such extensive use, antimicrobial resistance began to be reported as early as the 1940s, threatening the use of antimicrobials in medicine. Bans of antimicrobials as growth promoters did not appear until the mid 1980s.¹⁷

Considering the examples of late lessons from early warnings for a number of previous medical technologies, it is important to address risks of such emerging technologies at an early stage of development, which is where we find nanorobots today. In this paper, we present an early review of environmental and health risks related to nanorobots. First, we provide a brief overview of the progress in nanorobot developments, including which types of nanorobots are under development and in which applications they are likely to first emerge. Second, we review existing knowledge about nanorobots in the context of environmental and health risks, identifying a set of potential hazards to be further studied and scrutinized. Third, we discuss how nanorobots, if proven risky to the environment or human health, might become regulated. Finally, recommendations for the further enquiry into potential risks of nanorobots are provided.

A precise definition of the term ‘nanorobot’ is currently lacking. It is here tentatively defined as *an individual nano-sized device able to perform a designated task*. The nanometer size (referring approximately to the 1–100 nm size range) follows naturally from the term nanorobot, while the ability to perform tasks is central to any (also a macro-sized) robot and sets nanorobots apart from conventional, passive nanomaterials. This definition is similar to the one provided by Gao and Wang¹³ for ‘nanomachine’: *a nanoscale device that performs a task*. The main difference is that our definition specifies

individual devices themselves as being nanorobots, rather than devices consisting of entire swarms of nano-sized robots. It should be noted that this definition excludes a range of micro-sized robots. A specific example is MagnetoSperm, aimed at imitating a sperm cell in movement.¹⁸ This device has a thickness, length and width of about 5.2, 320 and 42 μm , respectively, which is here considered too large for a nanorobot. Another excluded example is the artificial bacterial flagellum described by Qiu *et al.*,¹⁹ which is about 1.2 μm wide and 16 μm long. Also excluded are metal–organic–framework-based biomedical microrobots, called MOFBOTS, which are at least a few micrometers in diameter.²⁰ Naturally, the delimitation to the nanoscale also excludes larger micro-sized robots, such as the 400 \times 800 μm tumbling micromotor.²¹ The exclusion of such micro-sized devices in this review is done to enable a focus on nano-sized devices, which might potentially exhibit unique hazardous properties due to their small size.²² However, we have been rather inclusive in our selection, including for example robots of some hundred nanometers in at least one dimension, even though most contemporary definitions of nanomaterials have an upper size limit of 100 nm.²³

2. Technological development of nanorobots

The technological development of both micro- and nano-sized robots stretches from Richard Feynman’s famous talk ‘There’s Plenty of Room at the Bottom’ in 1959 until now, intertwined with the development of other technologies, such as magnetic control and nanostructure synthesis.²⁴ The first nano-sized robots were reported in 2005, including the nanocar made mainly by fullerene molecules.²⁵ During the 2010s, three main designs of nanorobots have emerged: helices (also called nanoswimmers), nanorods (also called nanoswimmers, nanomotors or, if longer, nanowires) and DNA nanorobots. These will be described in more detailed in Sections 2.1–2.3 and a non-exhaustive list of nanorobot examples is presented in Table 1. These three categories generally fulfill the tentative definition of nanorobot in section 1 by being both nano-sized and able to perform some tasks. However, sometimes the task to be performed has not yet been entirely realized for specific nanorobot designs, but there is generally an ambition to execute successful drug delivery or some similar task, like biological imaging. In general, it seems that the nanorods and DNA nanorobots have reached the furthest towards actually accomplishing drug delivery and similar tasks.

2.1 Helices

A number of robots with screw-like helix tails for movement have been developed, often resembling bacterial flagella or other biological entities. Most of them are rather to be categorized as micro-sized robots, including the above-mentioned MagnetoSperm and the MOFBOTS. However, there exist examples of helix-like robots that are approaching the



Table 1 Non-exhaustive list of examples of nanorobots. N.a. = not available

Nanorobot	Materials	Nanometer size	Propulsion/navigation	Reported speed ($\mu\text{m s}^{-1}$)	Reference
<i>Helices</i>					
Helical propeller	Glass, cobalt	200–300 nm wide	Magnetism	40	26
Artificial bacterial flagellum	Indium, gallium, arsenic, chromium, nickel, gold	200 nm wide	Magnetism	1.2 (average)	27
Repolymerized flagella-bound nanoparticles	Magnetic materials, bacterial flagella	Nanoparticles: 40–400 nm Flagella: 20 nm wide	Magnetism	2.5 (max)	29
<i>Nanorods</i>					
Drug-delivering nanorod	Gold, nickel, polymer	250 nm wide	Ultrasound/magnetism	50 (max)	31
Antibacterial nanorod	Gold, lysozyme	<300 nm wide	Ultrasound/bioreceptors	n.a.	32
Coated nanowire	Gold, graphene oxide, DNA	200 nm wide	Ultrasound/bioreceptors	n.a.	33
Enzyme-bound nanowire	Gold, nickel, polymer, asparaginase	ca. 500 nm wide	Ultrasound/magnetism	32 (average)	34
Light-driven nanorod	Gold, iron oxide	300 nm wide	Visible light/magnetism	33	37
Light-driven nanowire	Silicon, platinum	500–1000 wide	Visible or near-infrared light/morphology	5–35	38
Match-like nanorod	Silica, silver, silver chloride	<210 nm wide	UV light, silver chloride decomposition	4–14	39
Nanorod with flagellum tail	Gold, nickel, polymer	300–600 nm wide	Acoustic waves/magnetism	60 (max)	46
V-Shaped nanorod	Platinum	700 nm wide	Hydrogen peroxide decomposition	2–7	41
Nanofish	Gold, nickel, silver	200 nm wide	Magnetism	31 (max)	42
Two-armed nanoswimmer	Gold, nickel, silver	200 nm wide	Magnetism	39 (max)	43
Machine-learning optimized nanoparticles	Polystyrene, platinum	398 nm	Hydrogen peroxide decomposition	n.a.	44
2D nanoswimmer	Barium ferrite, platinum	4.6 nm wide	Hydrogen peroxide decomposition/magnetism	n.a.	45
<i>DNA nanorobots^a</i>					
DNA walker	DNA	n.a.	Bioreceptors	n.a.	49
Molecular spiders	Protein, DNA	n.a.	Bioreceptors	n.a.	50
I-switch	DNA	n.a.	Diffusion/bioreceptors	n.a.	51
Hexagonal barrel cage	DNA	35 × 35 × 45 nm	Diffusion/bioreceptors	n.a.	52
Tetrahedron DNA nanoparticle	DNA	8 × 10 nm	Diffusion/bioreceptors	n.a.	53
Icosahedral DNA nanocapsules	DNA	n.a.	Diffusion/bioreceptors	n.a.	55
Nanosheet/tubular nanorobot	DNA	90 × 50 × 2 nm	Diffusion/bioreceptors	n.a.	56

^a Note that since DNA nanorobots are well-defined molecules, their size in terms of nm is not always provided. Also, their speed is not reported to the same extent as for helices and nanorods.

nanometer size range. One example is the helical cobalt-covered glass propeller developed by Ghosh and Fischer,²⁶ which is 200–300 nm wide and 1–2 μm long (Fig. 1a). The propeller is meant to mimic a bacterial flagellum in terms of swimming behavior. Thanks to the magnetic cobalt layer, the propeller can be moved and navigated through magnetic fields (both backward and forward), reaching speeds of about 40 $\mu\text{m s}^{-1}$. The navigation control in a water-based solution is exemplified by ‘writing’ micrometer-scale letters and symbols, such as ‘R’, ‘@’ and ‘H’, using the propeller’s trajectory. Several propellers can be controlled simultaneously in this way.

Another helix design that might be considered a nanorobot is the artificial bacterial flagellum by Zhang *et al.*,²⁷ which can be 200 nm thick, 2.5 μm wide and 2.5

μm long (Fig. 1b). It consists of a magnetic chromium–nickel–gold head attached to an indium gallium arsenide–gallium arsenide–chromium tail. It is the tail that resembles a bacterial flagellum (both in size and shape) and the head enables magnetic propulsion and control. The artificial bacterial flagellum can swim at an average speed of 1.2 $\mu\text{m s}^{-1}$ under magnetic fields thanks to its magnetic head. Similar to the helical glass propeller, the artificial bacterial flagellum can move both forward, backward and turn in water-based media depending on the shape of the helix and the direction of the magnetic field. It was furthermore shown that the artificial bacterial flagellum can be controlled into pushing and rotating polystyrene microparticles.



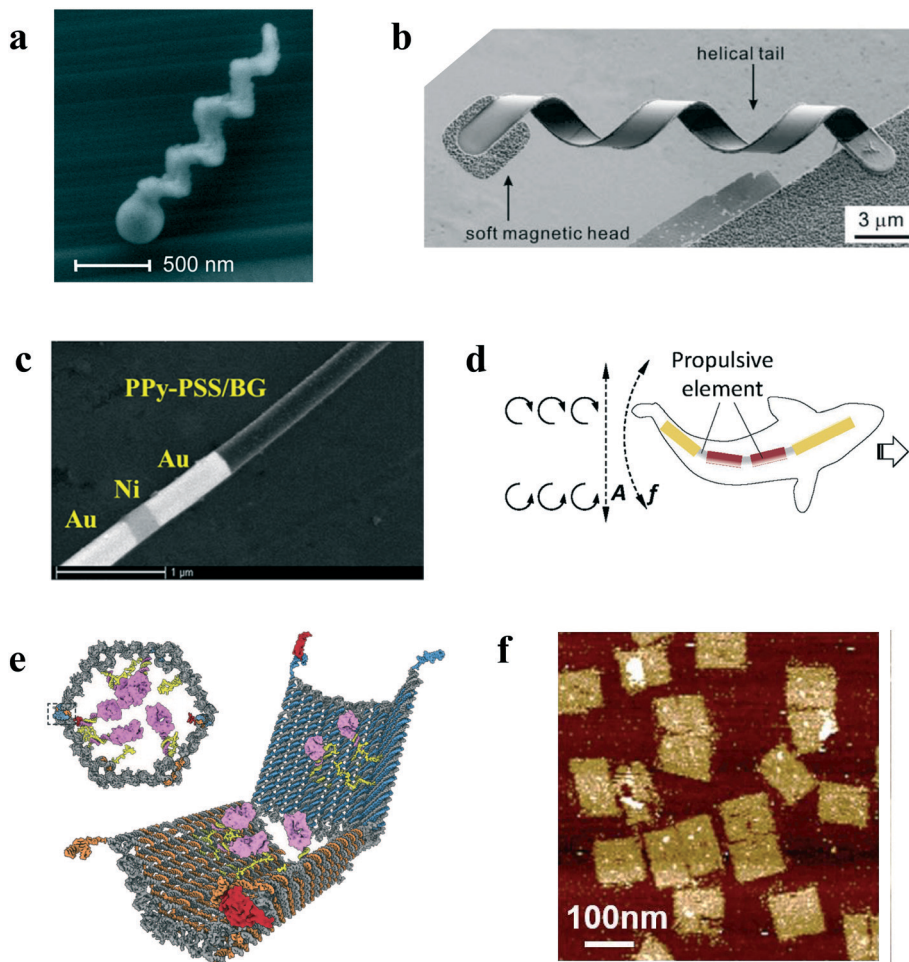


Fig. 1 Examples of nanorobots. (a) 200–300 nm glass propeller. Reprinted (adapted) with permission from Ghosh and Fischer.²⁶ Copyright (2009) American Chemical Society. (b) 200 nm thick artificial bacterial flagellum made by a chromium–nickel–gold head and an indium gallium arsenide–gallium arsenide–chromium tail. Reprinted (adapted) with permission from Zhang *et al.*⁵⁷ Copyright (2009) American Chemical Society. (c) Gold (Au)–nickel (Ni)–gold (Au)–polymer (PPyPSS) nanorod loaded with an antiseptic drug (brilliant green, BG). Reprinted (adapted) with permission from Garcia-Gradilla *et al.*³¹ Copyright (2013) American Chemical Society. (d) Fish-like 200 nm wide nanorod consisting of gold–nickel–nickel–gold segments with three flexible silver hinges linking the segments. Reprinted (adapted) with permission from Li *et al.*⁴² Copyright (2016) Wiley-VCH Verlag GmbH & Co. (e) Hexagonal cage-like DNA robot able to transport payloads (in pink). Reprinted (adapted) with permission from Douglas *et al.*⁵² Copyright (2012) Science. (f) DNA nanorobots consisting of 90 nm × 50 nm × 2 nm sheets able to fold into tubular drug carriers. Reprinted (adapted) with permission from Li *et al.*⁵⁶ Copyright (2018) Nature Publishing Group.

In addition to these two examples of nanorobot helices, several similar helix-like nanorobots have been developed, where most are micro-sized but some dwell somewhere in the borderland between the nano- and micro-size ranges.²⁸ There are also attempts to use actual bacterial flagella by de-polymerizing them into flagellin proteins using heating, then repolymerizing them back into flagella and attaching them to magnetic particles 40–400 nm in size.²⁹ Multiple flagella can be attached to the same nanoparticle, but a functional group that can bind to the nanoparticles must be introduced during the repolymerization. The repolymerized flagella have outer diameters of about 20 nm, lengths of 5–10 μm and can obtain different shapes (normal, curly and coiled) depending on if ethylene glycol or dimethyl sulfoxide is added. The idea is that reconfiguring its geometry might be beneficial when navigating through heterogenous biological environments (such as the human body) compared to

movement in pure water. By applying magnetic fields, the nanoparticle–flagella clusters can swim at velocities up to 2.5 $\mu\text{m s}^{-1}$. An advantage of this helical nanorobot design is that the specific nanoparticles can be changed for different purposes while maintaining the repolymerized flagella for movement. However, for both propulsion and navigation purposes, the nanoparticles must be magnetic.

2.2 Nanorods

The nanorods typically consist of cylindrical rods with different metal segments,³⁰ although different shapes are also used for the same purpose. A particularly notable example from a medical point of view is the 250 nm wide and 1800 nm long rod with gold–nickel–gold segments developed by Garcia-Gradilla *et al.*³¹ These nanorods move due to ultrasound waves



and can move in serum at about $50 \mu\text{m s}^{-1}$ and, albeit at lower speed, in saliva (about $10 \mu\text{m s}^{-1}$). Thanks to the magnetic properties of nickel, such nanorods can be steered along predetermined trajectories. For example, the developers made it 'write' the letters 'U', 'C', 'S' and 'D' with its trajectory. The nanorod can be functionalized, making it carry drug cargoes. An important potential application of the gold–nickel–gold nanorod was shown by functionalizing it with a polypyrrole–polystyrene sulfonate segment. This organic segment can bind to the antiseptic drug brilliant green (Fig. 1c) and deliver this drug to designated destinations. The drug can then become released due to changes in pH.

Another potential medical use of nanorods was demonstrated by Kiristi *et al.*³² They used ultrasound-powered porous gold nanorods less than 300 nm wide and functionalized them with the bactericidal substance lysozyme, which can kill both Gram-negative and Gram-positive bacteria. They showed that the lysozyme-functionalized gold nanorods could kill up to about 80% of the Gram-positive bacteria *M. lysodeikticus* in a sample within a few minutes. The number of nanorods influenced the death rate, with about 5000 being required for a rate of 80%. Glycosidic bonds in the cell walls of bacteria act as bioreceptors for the lysozyme. Although there was no navigation control for this nanorobot, the movement in itself greatly increased the lysozyme–bacteria interactions and thus also the bacterial killing capacity *versus* pure lysozyme only.

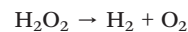
Several nanorod designs aim at cancer detection and treatment. MicroRNAs (miRNA) are small RNA strands, some of which can be associated with diseases such as cancer and diabetes. In the work by Esteban-Fernández de Ávila *et al.*,³³ graphene oxide-covered gold nanowires 200 nm wide were used to detect miRNA. The ultrasound-propelled nanowires can penetrate cancer cells where DNA strands bound to the graphene oxide surface detach and bind to the miRNA. This causes the graphene oxide–gold nanowires to send out a fluorescence signal, thereby detecting the presence of a specific miRNA and thus possibly also a cancer cell.

Moving beyond detecting cancer and towards treating it, Uygun *et al.*³⁴ used gold–nickel–gold–polymer nanowires as an effective anti-cancer agent. These nanowires were propelled by ultrasound at an average speed of $32 \mu\text{m s}^{-1}$ in human serum and magnetically guided thanks to the nickel. They furthermore had asparaginase enzymes bound to the polymer segment, which can deplete cancer cells of the essential amino acid asparagine and thereby inhibit their growth. A 92% inhibition of lymphoma cancer cells was observed, compared to only 17% inhibition for free, non-nanowire-bound asparagine.

While the chemical (*e.g.* hydrogen decomposition), magnetic, acoustic (*e.g.* ultrasound) and biological (*e.g.* attaching bacterial flagella) have conventionally been the main mechanisms of propulsion and navigation for nanorods,³⁵ recent studies have also reported using light for nanorod propulsion.³⁶ An example is 300 nm wide gold–iron oxide nanorods, which can be powered by visible light in diluted hydrogen peroxide, reaching speeds of $33 \mu\text{m s}^{-1}$ due to

hydrogen peroxide decomposition at the iron oxide end.³⁷ Due to the superparamagnetic iron, magnets can be used to steer the rods in designated trajectories. Similarly, platinum nanoparticle-coated silicon nanowires in a quinone solution can achieve propulsion if irradiated with light at the visible or near-infrared spectrum.³⁸ Speeds of about $5\text{--}35 \mu\text{m s}^{-1}$ could then be achieved depending on the power intensity of the light. Changing the end-surface morphology of the nanowire can result in different propulsion patterns, from linear to circular. A final example of light-propelled nanorods is the <210 nm wide match-like silver–silica nanorod with a silver chloride tail.³⁹ The nanorods were dispersed in water and irradiated with UV light, inducing a photocatalytic decomposition of the silver chloride tail. Depending on the length of the match-like nanorod, speeds of $4\text{--}14 \mu\text{m s}^{-1}$ can be achieved.

In addition to these examples of nanorods, often with intended or even realized medical applications, several other nanorod designs exist, often being about 200–300 nm wide and including the elements nickel, gold and/or platinum as segments.⁴⁰ However, nanorobots with shapes other than purely cylindrical but otherwise similar to nanorods have also been developed. One example is a 700 nm wide and 4000–4500 nm long V-shaped platinum nanorod.⁴¹ It can move in a hydrogen peroxide medium through the decomposition of hydrogen peroxide occurring predominantly at the non-pointy end of the V shape:



This creates a flow of oxygen, which propels the V-shaped nanorod forward at speeds of $2\text{--}7 \mu\text{m s}^{-1}$ depending on the hydrogen peroxide concentration. The directional movement of this nanorobot is thus achieved through its V shape, since the hydrogen peroxide decomposition occurs predominantly at one end of the robot. However, the nanorobot could only rotate in micrometer-wide circles and no navigational control was imposed. Another example of a nanorod consisting not only of a single cylinder shape is the 200 nm wide and 4800 nm long nanofish, consisting of several cylindrical segments joined together: a gold segment as head, two nickel segments as the body, a gold segment as the caudal fin and three flexible porous silver hinges linking the other segments (Fig. 1d).⁴² The nanofish can be propelled magnetically due to the nickel segments, swims by waving its tail in a manner similar to actual fish. The nickel segments in the fish body continuously align themselves with the orientation of the magnetic field, making the gold segment in the caudal (tail) fin exhibit undulatory motion, enabling the nanofish to reach a speed of about $31 \mu\text{m s}^{-1}$. A third example of a non-cylindrical-only nanorod type of nanorobot is a magnetically-controlled two-armed nanoswimmer consisting of 200 nm wide nickel–gold–nickel segments with silver hinges in between.⁴³ This nanorobot thus have two nickel arms doing the swimming and a gold body. In oscillating magnetic fields, this two-armed swimmer can reach a speed of about $39 \mu\text{m s}^{-1}$ in *e.g.* seawater. However, the speed achieved in serum was notably slower – approximately $10 \mu\text{m}$



s^{-1} . To our knowledge, the use of these non-cylindrical-only nanorods for medical purposes, such as drug delivery, have not yet been proven by experiments, although several of the authors express an ambition towards medical applications, arguing for example that their nanorobot designs and/or propulsion strategies are feasible in human bodies.

Adopting a structured approach to nanorobot design, Zeng *et al.*⁴⁴ used machine learning to design optimal micro-/nanorobots for the purpose of catalytic water cleaning, assuming hydrogen peroxide decomposition propulsion. Different aspect ratios, shapes, catalytic materials and other parameters were tested. The results suggested a spherical 398 nm platinum-coated polystyrene nanoparticle, which was then synthesized and successfully used for decomposing methyl blue dyes. In addition, two-dimensional (2D) nanoswimmers were designed by the same group, consisting of 4.6 nm thick platinum-coated barium ferrite platelets.⁴⁵ Again, hydrogen peroxide decomposition was the propulsion mechanism, and steering was enabled by magnetic control of the iron in the barium ferrite. The 2D nanoswimmers showed excellent performance in catalytically removing stains on cloth. These two non-cylindrical nanorobots are thus clearly intended for remediation and cleaning rather than for medicine.

Finally, constituting some sort of a hybrid between nanorods and helices is a nanorobot consisting of a 300–600 nm wide rod with nickel-gold segments and an attached polypyrrole tail.⁴⁶ When exposed to acoustic waves, the polypyrrole tail begins to oscillate and propels the nanorod forward at speeds up to $60 \mu\text{m s}^{-1}$. Although this nanorobot does not consist of a helical structure, and is thus classified as a nanorod, the polypyrrole tail acts much like an artificial flagellum in a similar way as the tails of some helical nanorobots (see section 2.1).

2.3 DNA nanorobots

DNA nanorobots consist of deoxyribonucleic acid molecules, thus using DNA as construction material for nano-sized devices.^{11,47} Sometimes, they are based on DNA origami, where DNA molecules are folded to create patterns and shapes.⁴⁸ An example of such a nanorobot is the DNA walker developed by Gu *et al.*,⁴⁹ which consists of a trigonal arrangement of double helices, resembling a symmetrical three-legged wheel with ‘feet’ that act like ligands. These DNA ‘feet’ can bind to a larger DNA origami sheet ‘landscape’, across which the DNA walker can ‘walk’ by rotating 120° and binding to a new bioreceptor in each step. It can deliver cargo across the DNA sheet landscape to a designated site, illustrated by the DNA walker delivering several 5 nm gold nanoparticles in the study. Other structures able to walk across a DNA origami ‘landscape’ are the molecular spiders, which consist of protein bodies with three DNA ‘legs’ and a fourth capture ‘leg’, specifically made by so-called DNA enzymes.⁵⁰ Due to the detailed design of the DNA origami ‘landscape’, the molecular spiders can ‘walk’ across the landscape as the legs dissociate from one site and

reattach to a new site. Since the spiders have three legs, complete dissociation is hindered as dissociated legs are held in place due to the binding of the two other legs and quickly reattach. The spiders can be made to follow pre-designed one-dimensional tracks in the ‘landscape’ and even execute commands like ‘turn’ and ‘stop’. The capture leg is used to capture the spiders from solution and place them at the starting position. Although the DNA walkers and molecular spiders have impressive programmability, it should be noted that their controlled movement and navigation seems to be limited to pre-designed DNA ‘landscapes’.

Moving away from such landscapes, DNA nanorobots have also been used in the *in vivo* environments of living organism. An example of such a device is a DNA nanorobot called the I-switch, which consists of three DNA strands.⁵¹ The I-switch can change shape depending on pH and the two shapes emit light of different wavelength when the nanorobot is tagged with a fluorescent molecule. This property can be used for tempo-spatial mapping of pH changes in living organisms, as shown for both wild type and mutant nematodes (*C. elegans*). Fluorescence-tagged I-switches were translocated to certain nematode cells and taken up through receptor-mediated endocytosis so that pH changes in the cells could be tempo-spatially mapped. Since many phenomena in cells, including neurodegeneration and spermatogenesis, are modulated based on changes in pH in the range where the I-switch is sensitive to changing shape, there might be several mapping applications for this DNA nanorobot.

An example of a DNA nanorobot with potential medical implications is an origami-based hexagonal barrel-shaped cage-like robot with dimensions of $35 \text{ nm} \times 35 \text{ nm} \times 45 \text{ nm}$ (Fig. 1e).⁵² The cage can be loaded with various materials, such as gold nanoparticles and in particular biologically active payloads, such as antibody fragments. The cage has ‘locks’ that can be ‘unlocked’ by binding to protein receptor ‘keys’. This causes the DNA nanorobot to undergo a drastic reconfiguration that releases the payloads. The release of payload was shown in human cells, such as leukaemia cells and lymphocytes. Similar delivery of therapeutics *in vivo* was achieved by tetrahedral DNA nanoparticles.⁵³ The tetrahedral is made by six self-assembling DNA strands, to which six strands of a particular type of RNA, called siRNA, were bound. The siRNA can be used to silence target genes in tumors. The application was demonstrated by delivering the tetrahedral-bound siRNA to target tumors in nude mice through injection in the tail vein. Another similar DNA nanorobot is the cage-like icosahedral DNA nanocapsule which can be used to encapsulate biomacromolecules.⁵⁴ The nanocapsule can target specific cells and deliver the molecules to the cytosol.⁵⁵ The release of the cargo is controlled by photoirradiation. The application was illustrated *in vivo* by delivering a neurosteroid, which can promote neurogenesis and neuron survival, in the nematode *C. elegans*. A final example of a DNA nanorobot is a $90 \text{ nm} \times 50 \text{ nm} \times 2 \text{ nm}$ DNA sheet (Fig. 1f) that can fold into a tubular nanorobot.⁵⁶ The sheets can bind certain cargos while open



and then and fold into a tube, thus encapsulating the cargo. Furthermore, the nanorobot can be functionalized with DNA ligands on the outside of the tube, which can bind to a receptor protein specifically expressed in tumor cells on the inside of blood vessels and unfold. This way, the tubular nanorobot was intravenously injected and able to transport thrombin to target blood vessel tumors in mice, where the nanorobot unfolded to release its cargo. Since thrombin is a molecule able to kill cancer cells in blood vessels, this leads to tumor cell necrosis and tumor growth inhibition.

3. Potential nanorobot hazards

Several of the nanorobot designs described in section 2 offer the promise of significant health-related benefits, such as improved cancer therapy. However, considering what can be learned from previous late lessons with promising technologies offering great societal benefits, such as the X-rays and antimicrobials discussed in section 1, risks can outweigh the benefits for some applications of a technology. So far, only a few references to environmental and health risks can be found in the literature about nanorobots. Kostarelos⁵⁸ wrote briefly about the safety of nanorobots, commenting that nanorobots “will need to be toxicologically inert, degradable or expelled from the body”. One might note that this mainly refers to human toxicity and not subsequent environmental effects that might occur after the nanorobots have become expelled from bodies. Gao and Wang¹³ wrote about the use of nanorobots (mainly nanorods) for environmental sensing, monitoring and remediation. They comment that “the potential toxicity of micro/nano-scale motors needs to be evaluated to prevent potential adverse environmental impacts”. However, they do not provide any specific recommendations on how that could be accomplished, despite envisioning wide-spread use of nanorobots in the environment. Surana *et al.*⁵⁹ did a study on DNA nanorobots and their compatibility with the immune system of higher organisms. They comment that foreign, ‘non-self’ DNA from other organisms can be harmful and therefore immunogenic, since they trigger the immune system: “Even though DNA is a natural biopolymer, when present at the wrong place at the wrong time it can elicit a strong inflammatory reaction”. Therefore, they asserted that it is important to consider the various cellular and systemic responses that such DNA architectures might elicit, which are likely to be species-specific. Such considerations have a dual purpose: it is both to keep the organism in questions safe from the DNA nanorobot but also to ensure the proper medical function of the DNA nanorobot in cells. Again, the focus is on human toxicological responses rather than on environmental toxicity.

Some consideration of safety can be found in studies describing DNA nanorobots. The developers of the I-switch noted that the nematodes injected where “viable and healthy”, indicating that the I-switch is non-toxic to nematodes given the applied concentrations.⁵¹ In the study by Li *et al.*⁵⁶ about the DNA nanosheet/tubular nanorobot, an assessment of the safety of the nanorobot was conducted. It was noted that the

nanorobots did not elicit any thrombi or increased blood coagulation in non-tumor-bearing mice at relevant concentrations. In addition, no immunological or cytotoxic responses were shown. They also did not cause any thrombi or blood coagulation in Bama miniature pigs, which is an animal similar to humans in anatomy and physiology. Although these results provide an early indication that such nanorobots might be safe, they are also limited to human toxicity impacts.

To these considerations of nanorobot risks in the previous literature, we might add a number of potential hazards. That foreign DNA can elicit immunologic and inflammatory responses has been noted above. Other materials used in contemporary nanorobot designs (see *e.g.* Table 1) might also potentially have hazardous properties that warrant further investigations. For example, the nickel used in order to magnetically control the propulsion of several helices and nanorods is known to be allergenic, carcinogenic (though not in pure metallic form), toxic at high doses and in certain forms, as well as teratogenic at high doses.⁶⁰ Allergic reactions have already been seen for people working with nano-sized nickel powder.⁶¹ The silver sometimes used for making hinges in designs with several connected nanorods is also known to be toxic to organisms in the environment – both in nano-form and when dissolved into silver ions.^{62,63} However, silver is not toxic to humans. High silver intake results in discoloration of the skin and internal organs (argyria and argyrosis, respectively), both which do not seem to bring any negative health effects.⁶² In addition, the UV light used for propulsion in some nanorod designs is known to be able to cause skin damage and, in the worst case, skin cancer.

Foreign DNA, nickel, silver and UV light are all established hazards. Whether their use in specific nanorobot applications constitute risks remains to be investigated. Novel hazards associated with nanorobots might be related to the control of nanorobot propulsion and navigation – whether by chemical propulsion, magnetic fields, sound waves, bioreceptor binding and/or light – potentially making the nanorobots travel to places in the human body and elsewhere where they are not supposed to. Should loss of propulsion control or targeting of an erroneous site occur, hazardous drugs might be delivered to healthy cells. An erroneous targeting might cause high concentrations locally, so that a small number of nanorobots potentially causes much harm.

Besides potential hazards, an additional aspect of risk is whether organisms will become exposed to the potential hazard. Whereas nanoparticles have typically been perceived as extrasomatic risks, released to the environment and subsequently taken up by organisms,⁶⁴ the mainly medical applications envisioned for nanorobots mean that exposure and uptake to humans might be inherent in the use of nanorobots rather than unintentional. Environmental exposure might then potentially occur subsequent to excretion or discarding of the nanorobots. In addition, the use of nanorobots for environmental remediation also seems to imply a direct exposure to organisms in the environment,¹³ in that sense being similar to pesticides



applied to agricultural land. The probability of nanorobot exposure to relevant organisms might thus be high for these two promising applications.

4. Regulating nanorobots

As in the early development of X-rays and antimicrobials, there is currently no regulation targeting the use of nanorobots specifically. Meeting the current approval requirements for medical products and devices is arguably one of most lengthy, thorough and expensive regulatory processes around, with various phases of clinical testing, safety and benefit assessment. However, regulations in the EU and elsewhere have still been criticized for being insufficient when it comes to more complex drugs.⁶⁵ It even remains unclear whether nanorobots are to be considered a medical device or a medicinal product, for which different sets of regulations apply in the EU – the Regulation of Medical Devices and Medicinal Products Directive, respectively. Currently, the ‘mechanism of action’ is key to decide whether a product should be regulated as one or the other. These can be pharmacological, immunological or metabolic means, which is why the categorization of nanorobots would depend on if they use complex mechanisms of action combining mechanical, chemical, pharmacological and immunological properties, as well as if they have both diagnostic and therapeutic functions.⁶⁶

One of few regulations that includes specific considerations for nanomaterials is the EU Regulation on Medical Devices (Regulation (EU) 2017/745),⁶⁷ where nanomaterials are defined according to the European Commission’s recommendation: “a natural, incidental or manufactured material containing particles in an unbound state or as an aggregate or as an agglomerate and where, for 50% or more of the particles in the number size distribution, one or more external dimensions is in the size range 1–100 nm”.⁶⁸ This definition, currently under revision, was clearly developed with nanoparticles in mind (considering the reference to particle size distribution) and not with a focus on active nanomaterials. An annex of the Medical Devices Regulation calls for “special attention” to be given to nanomaterials without further specification, and again focuses on nanoparticles: “devices shall be designed and manufactured in such a way as to reduce as far as possible the risks linked to the size and the properties of particles which are or can be released into the patient’s or user’s body, unless they come into contact with intact skin only.” If there is a high or medium potential for internal exposure, devices incorporating or consisting of nanomaterial should also be classified as ‘Class III’. This constitutes the highest risk classification in the EU and is reserved for the most critical devices, for which explicit prior authorization with regard to conformity with current standards and rules is required for them to be placed on the market. However, these rules for classification of medical devices and conformity assessment procedures were adopted in 1993 – a decade before the more widespread use of nanotechnology. Current regulations thus seldom include nanomaterials, and when they do, the focus is on

nanoparticles. Already more than ten years ago, when nanorobots were hardly developed, the European Medicines Agency⁶⁹ argued that appropriate expertise will need to be mobilized for the evaluation of the quality, safety, efficacy and risk management of novel applications of nanotechnology, such as nanostructures allowing transport across biological barriers, remote control of nanoprobe, and multifunctional chemical structures for drug delivery and targeting of disease. Back then, such novel applications of nanotechnology did not exist, but current developments of nanorobots are increasingly making such applications possible.

5. Recommendations

The main applications envisioned for nanorobots are such that they might potentially become administered directly to the human body or the environment (section 3). Such applications with potential for exposure, akin to those of pharmaceutical and pesticide applications, warrant consideration into the risks related to nanorobots. We identified two main potential hazards related to nanorobots at this early stage: (i) the use of conventional hazards, such as hazardous materials and UV light, as well as (ii) the loss of propulsion and navigation control (section 3). Furthermore, we note a lack of nano-specific regulation, making it uncertain whether current regulation will be able to identify and regulate nanorobot hazards at an early stage of development (section 4). In order to address this situation, we provide three recommendations for future research and action. The recommendations are based on three lessons learned from failing to respond to early warnings in the past,¹⁴ which seem particularly relevant to the discussion about nanorobot risks: (i) acknowledge and respond to ignorance, uncertainty and risk in technology appraisal, (ii) ensure use of ‘lay’ knowledge, as well as specialist expertise, and (iii) systematically scrutinize claimed benefits and risks. Following the three recommendations would allow for making the most of nanorobots while avoiding that their use later turns out to cause harm to the environment and/or human health.

The first recommendation, based on lessons (i) and (iii), is to conduct studies of the environmental and human health risks of different nanorobot designs before they are in widespread use, moving away from the view that nanoparticles are the only aspects of nanotechnology for which risk assessment and regulation are needed. With nanoparticles, discussions about their risks started early in the development of nanotechnology. We have since then learned how important it is that sufficient funding of risk-related research is provided and that studies are initiated early on in order to map different risks. Although it is currently unknown whether nanorobots constitute a potential risk to human health and the environment, it is possible to start processes where this can be investigated. At the moment, risk assessments of medical devices and medicinal products are largely focused on establishing whether a drug or device is safe to use under a specific set of conditions using pre-defined test methods, which means that novel risks



Table 2 Ten questions recommended to be addressed in risk-related studies of nanorobots

1. What is the toxicity of nanorobots and their constituents to humans and other organisms?
2. Are nanorobots more hazardous than previous generations of passive nanomaterials?
3. How many nanorobots are expected to be produced and used in the future?
4. What is the likely future exposure of nanorobots to humans and organisms in the environment?
5. In which ways can the propulsion and navigation of different nanorobots be obscured?
6. How can existing regulations be adapted to cover potential risks of active nanomaterials such as nanorobots?
7. How can nanorobots be designed to be safe?
8. How can the benefits of nanorobots be quantified and compared to the potential risks?
9. What is peoples' risk perception of nanorobots?
10. What are the main societal concerns related to nanorobots?

will probably come as surprises under the current regulatory framework. It would therefore be wise to conduct broader studies of the potential risks of nanorobots, considering different potential hazards and other risk-related aspects. Table 2 provides a non-exhaustive list of ten questions which we recommend be addressed in such risk-related studies of nanorobots, including the consideration of the potential hazards outlined in section 3. Among these, question 3 about the future production and use of nanorobots is fundamental. Currently, nanorobots have only begun to be tested at laboratory scale for applications such as medicine and environmental remediation. Their future production and use will depend on the technical performance of nanorobots in these applications. In medical applications specifically, an important prerequisite for successful use is that the nanorobots can evade the immune systems of the organisms.⁷⁰ For DNA nanorobots in particular, this can be challenging since foreign DNA is immunogenic,⁵⁹ although conducted tests for the tetrahedron DNA nanoparticles⁵³ and nanosheet/tubular DNA nanorobot⁵⁶ did not indicate any immunogenic response in mice. Another prerequisite for future nanorobot use that convenient, large-scale fabrication methods are achieved. Without a high-enough production and use of nanorobots, any associated risks will remain low. However, it should be noted that all technologies are sparsely used in the earliest beginning of their development, in their so-called embryonic phase.⁷¹ Therefore, an initially low production and use should not be taken as evidence of future low production and use. Detailed monitoring of production volumes, along with technology forecasting and scenario analyses, are recommended for addressing question 3.

The second recommendation, based on lesson (ii), is that policy-makers and regulators should reach out to relevant expert stakeholders and initiate dialogue about how, when and why nanorobots might be used and address some of the issues that we know from discussions about nanoparticles are likely to be contentious for the future of nanorobots. Such issues include the discussion about nanomaterial and nanoparticle definitions, which are on-going but have not yet resulted in consensus.²³ Over time, stakeholder positions seem to have become increasingly entrenched with the emergence of increasing evidence indicating that nanoparticles might be associated with harmful effects on the environment and human health.⁷² There is no need to

wait with discussions about definitions until early indications of harm emerges. Discussions about a regulatory relevant definition of nanorobots should start now when the stakes are not yet so high for stakeholders involved. The tentative definition of nanorobots provided in section 1 of this paper provides a starting point for such discussions.

The third recommendation, also based on lesson (ii), is the initiation of broader public dialogue about risks and benefits of nanorobots, as well as how regulatory measures can be implemented in order to maximize the benefits of nanorobots while minimizing potential risks. For nanotechnology in general, there was an early effort already in 2004 to explore of the general public's attitudes towards nanotechnologies.⁷³ Many of the identified areas of concern are still highly relevant for nanotechnologies in general and nanorobots in particular. For instance, the public raised questions about the purpose and controllability of nanotechnologies, whether health and environmental considerations had been adequately addressed, whether existing regulation was up to the task, and whether lessons from the past had been learned. For nanorobots, we recommend reengaging in a dialogue with the public, listening to their concerns, and ensuring these concerns are addressed up-front.

Conflicts of interest

There are no conflicts to declare.

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