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Applications of guanine quartets in nanotechnology and chemical biology

Loic Stefan¹* and David Monchaud²*

Abstract | Guanine and related nucleobases such as guanosine, deoxyguanosine and isoguanosine are notable molecular tools for designing functional supramolecular assemblies. This popularity originates in their ability to self-assemble via a unique topological pluralism — as isolated nucleobases, discrete macrocyclic quartets and virtually infinite linear ribbons — that endows them with a considerable functional versatility. Many programmes have been launched to fine-tune the chemical properties of guanine derivatives, to make them usable under different experimental conditions, such as in organic or aqueous environments, and responsive to external stimuli, such as ionic strength, pH, light or temperature. These strategies aim to translate the chemical information encoded in a basic guanine unit into programmable, higher-order supramolecular architectures. Spectacular results have been recently obtained in various chemical fields, from supramolecular chemistry to chemical biology, from soft matter to catalysis. In this Review, we detail these advances and demonstrate how these multidisciplinary investigations cast a bright light on the diversity that guanines, synthetic guanines and related nucleobases uniquely offer in terms of both structure and function.

Guanines and related nucleobases such as guanosine, deoxyguanosine, isoguanosine and so on (simply referred to as G hereafter) are fascinating chemical tools, owing to their ability to create programmable supramolecular networks through their self-association to form G-quartets or G-ribbons, i.e. macrocyclic arrangements of four guanines or virtually infinite linear chains of guanines, respectively¹ (BOX 1). These self-assembled structures are stabilized by the formation of H-bonds at both the Watson–Crick² and Hoogsteen³ faces, making guanine a highly versatile tool for the assembly of supramolecules, with steadily increasing applications in both nanotechnology and chemical biology.

The history of guanine

The discovery of the unique properties of guanines can be traced back to the turn of the twentieth century (FIG. 1). Stubborn scientists were determined to elucidate the chemical nature of nucleic acids⁴, first isolated in 1871 by Miescher⁵. A laborious isolation of guanylic acids from bovine pancreas was achieved by Hammarsten in 1894 (REF.⁶), but uncertainties remained as to the monomeric or polymeric nature of this novel, non-protein natural substance. Independent efforts by Levene and Jacobs⁷ (who coined the term tetranucleotide hypothesis) and Bang⁸ towards the end of the first decade of the 1900s provided ever purer guanylic acid and, in turn, solid evidence supporting the monomolecular nature of guanylic acid. They noted, as did others, the tendency of concentrated solutions to 'gelatinize' in a vivid manner, describing guanylic hydrogels as "an amorphous, somewhat gelatinous precipitate (that) turns into a semiliquid jelly"⁹. The precise structural nature of the viscous guanylic hydrogel linchpin was eventually unveiled by Davies and colleagues in 1962 (REF.¹⁰), in a report describing the very first X-ray crystal structure of a G-quartet. It seems unlikely that even the most clever of scientists would have envisioned at that time the tremendous potential of this 'semiliquid jelly' to, a century later, become a key player in supramolecular chemistry, genetics, chemical biology and catalysis.

Native versus synthetic G-quartets

The current fascination with G-quartets undoubtedly originates from their involvement in the formation of higher-order nucleic acid structures, known as G-quadruplexes. The discovery of G-quadruplexes led to a paradigm shift in genetics because they "contribute spatial and temporal dimensionalities to linear sequence information" (Maizels)¹¹. This paves the way for a new view of DNA in which it is not merely a barcode waiting to be scanned but actively controls its own expression according to a new — and not yet fully understood — dimensionality provided by local formation of quadruplexes. DNA then has a structurefunction relationship and joins RNA (for which this has long been accepted). In this Review, we do not aim to discuss at any length the roles of the G-quartets when

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Box 1 | H-bonded guanines

Guanine (and related guanosine, deoxyguanosine) shines by its hydrogen-bond (H-bond) donating/accepting capability, with three acceptor (A) and three donor (D) sites, which enables both Watson–Crick and Hoogsteen faces to participate in the creation of a broad array of H-bond networks¹⁵². The iconic guanine/cytosine (G≡C) base pair follows the rule described by Watson and Crick², involving three H-bonds on the sole guanine's Watson–Crick face. When both faces are implicated, the supramolecular assemblies depend on the experimental conditions: in the presence of cation (e.g. K⁺), the guanosine (G) quartet formation is privileged, given that alkali metals can neutralize the electrostatic repulsion of the inwardly pointing guanine's carbonyl groups within the quartet inner cavity^{1,20}. The stability of the macrocyclic edifice thus originates in an interplay between the formation of eight H-bonds, the cation-mediated electronic stabilization, along with the delocalization of the nucleobase electrons (resonance), either π -electrons or σ -electrons or both (the resonance-assisted hydrogen bonding (RAHB)¹⁵³ or the charge separation theory¹⁵⁴, respectively). In the absence of cations, G-ribbon formation is privileged: two types of ribbon are possible (type A and type B), which differ from the orientation of the guanine's molecular dipole (arising from the highly polarized aromatic surface of the nucleobase, yellow arrows)^{1,155}. The nature of the solvent will favour ribbon with or without global dipole moment.



embedded in G-quadruplexes or the prevalence of quadruplex structures in the human genome, transcriptome and ncRNAome (known as G4ome)¹². These topics have been frequently covered in a number of other authoritative reviews^{13–17} and readers are directed there. Here, we focus on both natural and non-natural G-quartets or synthetic G-quartets (SQ) — used as molecular platforms for nanobiotechnological purposes.

The field of G-based molecular tools was launched in 1995 independently by Davis et al.¹⁸ and Gottarelli et al.¹⁹, who demonstrated that lipophilic guanine (LipoG) derivatives (5'-*tert*-butyldimethylsilyl-2', 3'-isopropylidene-isoguanosine (**1**) and 3',5'-didecanoyl-2'-deoxyguanosine (**2**), respectively; FIG. 2a) fold into stable, cation-templated SQ in organic media. In-depth NMR investigations performed with both systems indicated that potassium ions trigger the formation of higher-order assemblies, $[(1)_4K^+(1)_4]$ and $[(2)_4K^+(2)_4]$, respectively, in which the potassium ion is sandwiched

between two layers of SQ (so-called octamers; FIG. 2b). Both studies demonstrated the specific ion binding of SQ-based systems, with a better chelation of potassium (K⁺, 1.33 Å) versus the smaller sodium (Na⁺, 0.98 Å) (and no chelation for the even smaller lithium (Li⁺, 0.68 Å)), confirming the privileged shape and charge recognition of K⁺ by G-based suprastructures²⁰. Interestingly, Davis et al. soon demonstrated the importance of the chemical nature of the G-derivatives: while guanosines are selective for K⁺, isoguanosines selectively interact with larger cations such as caesium (Cs+, 1.67 Å) and radium (Ra2+, 1.48-1.70 Å), adapting the nature of its supramolecular assembly from octamer to decamer, $[(1)_5Cs^+(1)_5]^{21}$ and $[(1)_5 Ra^{2+}(1)_5]^{22}$, respectively (and, thus, from quartets to quintets). This opened the way to assemble a G-based toolbox in which to select G-derivatives in an application-driven manner (see below).

Efforts were, thus, subsequently invested to fine-tune the chemical structure of LipoGs to endow

G4ome

the G4ome comprises all G-quadruplex (G4)-forming sequences identified in the genome (G4-DNA) and both the coding and non-coding transcriptome (G4-RNA).

LipoG

lipophilic guanine derivatives aimed at being used in organic solvents, masking hydrophilic moieties (e.g. the hydroxyl groups of the guanosine's ribose) through the use of aliphatic protecting groups.



Fig. 1 | Timeline of the major steps in the discovery of DNA and G-quartets.

From the first observation by Pasteur and Virchow on cell division (discarding the notion of 'spontaneous generation'), the history of DNA is marked by many important breakthroughs (grey boxes), chief among them the isolation of DNA ('nuclein') by Miescher¹⁵⁶, the demonstration of its role as a repository of genetic information by Avery¹⁵⁷, the elucidation of its duplex structure by Watson and Crick² and the central dogma of biology heralded by Crick¹⁵⁸. Our understanding of G-quartets has developed alongside, while also being fully part of the history of DNA, with its own important steps (blue boxes): the isolation of guanylic acid by Hammarsten⁶, the description of guanylic hydrogel by Levene⁷ and Bang⁸, the elucidation of the guanosine (G)-quartet structure by Davies¹⁰, the first G-based devices by Davis¹⁸ and Gottarelli¹⁹, and the demonstration of the existence of G-quartet-based suprastructures referred to as G-quadruplexes in human cells by Balasubramanian^{159,160}.

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them with special properties. For instance, Rivera and colleagues designed monomeric guanines that self-assemble into SQ with an extended H-bond network (increasing from 8 to 12 H-bonds)²³ to improve their stability. They synthesized a LipoG derivative with an expanded Hoogsteen face, adding an acetylphenyl appendage on its C8-position (8-(3-acetylphenyl)-(3',5'-bis-isobutyryl)-2-deoxyguanosine (**3**); FIG. 2a). This allows for the formation of a synthetic G-quartet in which the amine moieties located in the outer side of the SO, and, thus, readily accessible to H-bond-donating groups²⁴, are involved in additional intramolecular interactions. In addition to higher stability, the resulting SQ display extended π -surfaces that result in the formation of K+-promoted hexadecameric columnar assemblies ($[(\mathbf{3})_{4}K^{+}(\mathbf{3})_{4}K^{+}(\mathbf{3})_{4}K^{+}(\mathbf{3})_{4}]$). These assemblies can be topologically switched to octameric assemblies by replacing K⁺ with strontium (Sr²⁺, $[(\mathbf{3})_{4}\mathrm{Sr}^{2+}(\mathbf{3})_{4}])^{25}$, in a reversible manner, on the basis of electrostatic consideration only (FIG. 2c). Spada and colleagues also exploited the strategic C8-modification of LipoGs to introduce a photo-switchable styryl moiety (3',5'-O-diacetyl-8-styryl-guanosine (4); FIG. 2a), which enables control over the K+-promoted formation of the $[(4)_{4}K^{+}(4)_{4}]$ octamers and their light-induced disassembly²⁶. Irradiation at $\lambda_{ex} = 365$ nm indeed triggers the *E*-to-*Z* conformational switch of the 8-styryl-guanosine, creating a steric clash that precludes π -stacking interactions between SQ planes and, thus, causes the supramolecular complex to disaggregate. Upon irradiation at $\lambda_{ex} = 254$ nm, the conformational switch is reversed (Z-to-E conversion) and the octamer restored, making the system fully reversible (no cycling wastes) and controllable (FIG. 2d). C8-modified LipoGs were also useful for gaining insights into the many external factors that influence the topological dynamism of resulting SQ. Using 8-bromo-2',3'-O-isopropylidene-5'-tert-butyldimethylsilyl-guanosine (5) (FIG. 2a), Meijer and colleagues confirmed the expected roles of solvent polarity and of the nature of the cation, but also highlighted a subtle yet significant influence of the Coulombic energy of cation-anion pairs (e.g. KCl, KBr or KI) used for templating SQ formation²⁷. Ion pair-separation energy, although largely negligible in water, is important in organic buffers, given that it may have an impact on the nature of the resulting supramolecular assemblies (octamers, dodecamers or hexadecamers). Brown and colleagues further substantiated these observations28, demonstrating that the topological pluralism of LipoG might also be subtly influenced by the method of investigation (in this case, the differences between solution and solid-state NMR), highlighting again the need to exercise caution when studying highly dynamic supramolecular systems. Anecdotally, 5 was a synthetic intermediate in a strategy implemented by Davis and colleague to take control over the topological diversity of G-based assemblies²⁹. Attempting to irreversibly freeze octamers via covalent crosslinking, alkene appendages were inserted on the C8 and N2 positions of 5. The resulting Ba²⁺-promoted octamer was then subjected to olefin metathesis, but this very elegant strategy proved unsuccessful.



Fig. 2 | LipoGs and their higher-order supramolecular assemblies. a | Monomeric lipophilic guanosines (LipoGs) investigated by Davis (1)¹⁸, Gottarelli (2)¹⁹, Rivera (3)²³, Spada (4)²⁶ and Meijer (5)²⁷ used as models to gain insight into the parameters that govern G-quartet formation. b, c | LipoGs (2, 3) were used to demonstrate the capacity of K⁺-promoted G-quartets to adopt higher-order suprastructures, such as octamers and hexamers. c, d | A variety of strategies were implemented to gain control over G-quartet topological pluralism (tetramer, octamer, hexadecamer), in a reversible manner, using either metallo-responsive or light-responsive LipoGs. SQ, synthetic G-quartets.



Fig. 3 | **Tetrameric lipophilic guanosines assemblies. a** | Guanosine (G)4-calix synthesis from (6)³² and spontaneous, Na⁺-promoted self-association leading to the formation of nanotubes based on the intermolecular synthetic G-quartets (SQ) thanks to the alternate (up–down–up–down) orientation of the 1,3-*alt*-calix[4]arene template. **b** | Template-assembled synthetic G-quartets (TASQ) synthesis from (**7**)³⁴ and representation of its 'open' (centre) and 'closed' (right) conformation, in which an intramolecular SQ is folded, owing to the all-up orientation of the cavitand template.

It was later revisited and applied efficiently for the synthesis of artificial ion transporters (see below).

A further possibility that enables greater control over the nature of the G-quartet-based suprastructures is to assemble multiple guanines within the same chemical scaffold. Lipophilic G-dimers and G-tetramers were designed to construct higher-order assemblies in a precise manner. The self-assembly of a covalently tethered C8-aryl G-dimer derived from 5 led to discrete quadruplex-type octamers (whose structure was confirmed by single-crystal X-ray diffraction). The resulting structures were found to be highly stable, even in protic (methanol, up to 100%) or polar aprotic (dimethyl sulfoxide, DMSO, up to 50%) solvents³⁰. To build supramolecular ion channels, Matile and colleagues synthesized G-dimers to assemble artificial ion channels but faced tremendous solubility issues, which likely originated from the use of non-glycosylated purine moieties³¹. Davis and colleagues successfully addressed

this issue, assembling four 5'-amino-2',3'-isopropylideneguanosines (6) around an alternating calix[4]arene scaffold³². The resulting G4-calix (FIG. 3a) orientates two guanines above and two guanines below the plane defined by the calixarene core (in an 'up-down-up-down' manner), so that the spontaneous, Na+-promoted assembly of multiple G4-calix moieties results in the formation of micrometre-long, SQ-based nanotubes. Sherman and colleague followed a similar approach, using the alkyne-azide cycloaddition to attach four 5'-azido-2', 3'-isopropylidene-guanosines (7) to a Cram-bowlshaped cavitand displaying four propargyloxy groups³³ to assemble lipophilic G-tetramers, referred to as TASQ (template-assembled synthetic G-quartets)³⁴. The all-up organization of the cavitand template makes TASQ the first example of discrete, intramolecular SQ. The presence of cations is not mandatory for the SQ formation; however, TASQ efficiently chelate Na⁺ or Sr²⁺, in a complex in which the quartet acts as a lid on top of a

c-di-GMP

G4-ligands

documented.

low-molecular-weight

3,3'-cyclic diguanylic acid

originally as a regulator of

discovered by Benziman and colleagues in 1987 (REF.¹⁵¹),

cellulose production in bacteria

and now known as a regulator

of virulence factor production and biofilm formation.

molecules designed to interact

through π -stacking interactions

external quartets, but groove

and loop binding has also been

specifically with DNA/RNA quadruplexes (G4s), mostly

with the G4's accessible

basket-like ion receptor (FIG. 3b), while larger cations such as Cs⁺ trigger TASQ dimerization [(TASQ)Cs⁺(TASQ)]³⁵.

Lipophilic versus water-soluble G-quartets

LipoGs have been instrumental in understanding the factors that govern G-quartet formation and stability, thanks to the broad variety of techniques implementable in organic solvents that are often more challenging to apply in water (see below). However, water-soluble guanines open up opportunities in terms of applications, notably for investigations in biological systems. From a structural point of view, the simplest hydrophilic monomer is guanosine monophosphate (GMP; FIG. 4). Early investigations on G-quartet formation were performed using concentrated GMP solutions (25 mg mL⁻¹)¹⁰. The structure of GMP-based, hydrophilic G-quartets was further investigated later on, through modern analytical techniques. For instance, using a combination of scanning electron microscopy (SEM) and NMR, Walmsley and colleagues demonstrated that Na⁺ indeed prompts GMP to form G-quartets [(GMP)₄Na⁺] that then spontaneously self-assemble into micrometre-length rods comprising thousands of stacked G-quartets. Meanwhile, K⁺ leads to the formation of both rod-like lattices (formed of stacked [(GMP)₄K⁺], similar to those formed with Na⁺) and twisting fibres comprising continuous helices of hydrogen-bonded guanines, with 15 Gs per turn³⁶ (FIG. 4a). The puckering of both the C2'-endo/C3'-endo sugars alternates along the strands and the right-handed helical twist of the suprastructure were confirmed by Wu et al., using a combination of 1D and 2D NMR spectroscopy techniques^{37,38}. GMP-dimers and GMP-tetramers have also been investigated. For example, the naturally occurring GMP-dimer c-di-GMP39, which is a central bacterial messenger, was not observed to fold into G-quartets or G-quadruplexes under the conditions found inside cells (300 mM K⁺ and 10 mM c-di-GMP). This observation discounts any possible biological role of c-di-GMP higher-order assemblies in the bacterial cell cycle. However, the presence of substoichiometric quantities of aromatic intercalators (such as proflavine³⁹ or thiazole orange⁴⁰; FIG. 4b) leads to the formation of highly stable c-di-GMP quadruplexes with a cage-like structure in which the aromatic inducer is intercalated between two layers of G-quartets. Sherman and colleagues scrutinized the structure of tetrameric GMP, following the lipophilic TASQ approach and exploiting the phosphate group of the 2'-deoxyguanosine-3'-monophosphate to link guanines to the cavitand template⁴¹ (FIG. 4c). As above, the presence of cations is not mandatory for the G-quartet formation of water-soluble TASQ; this new molecular tool was not exploited for cation chelation but as a DNA-mimicking platform to investigate the interactions between G-quartets and G4-ligands (vide infra).

Overall, the wealth of information collected over the past 25 years with both lipophilic and hydrophilic G-quartets has provided invaluable insights into the thermodynamic, dynamic and structural parameters that govern the assembly of G-quartet structures. The analytical techniques available to study these structures have become increasingly effective and accurate, making the area a playground for the development and application of cutting-edge technologies. From the pioneering crystallographic works of Gellert et al.¹⁰ that provided the very first details about the structure of a G-quartet, many different practical and theoretical techniques (often combined) have been applied to gain insights into the kinetics and thermodynamics of G-based assemblies. This includes other X-ray-based techniques (for example, small-angle X-ray scattering (SAXS))⁴², along with various microscopy techniques: atomic force microscopy (AFM) was applied to characterize the self-organization of GMP on mica⁴³, SEM was applied to investigate the folding parameters and global topology of micrometre-long Na₂(5'-GMP) assemblies³⁶ and scanning tunnelling microscopy (STM) was applied in combination with density functional theory (DFT) calculations to assess whether and how G-quartets can be assembled on surfaces (either Au(III)^{24,44} or highly oriented pyrolytic graphite (HOPG)45 surface) in solventless conditions, with simple or sophisticated guanine derivatives, in the absence or presence of Na⁺, K⁺ or Ca²⁺ cations. Spectroscopic techniques provided invaluable clues about G-based self-assemblies in solution; these include Fourier transform infrared (FTIR)⁴⁶, infrared multiple photon dissociation (IRMPD)47-49 and RAMAN50 spectroscopies, along with electronic and vibrational circular dichroism for instance (ECD⁵¹ and VCD⁵², respectively) combined with degenerate coupled oscillator (DCO) calculations⁵² and NMR^{53,54}, notably diffusion-ordered spectroscopy (DOSY) NMR, which were successfully implemented to characterize the cation-templated GMP aggregation in near-physiological conditions. Finally, theoretical models were provided via molecular dynamics (MD)55,56 and molecular mechanics (MM)57, notably for both describing cation-templated G-quartet formation and simulating in silico the movement (channelling) of cation (e.g. K⁺) between two G-quartet layers. Dispersion-corrected DFT was also found to be a quantum chemical computational method of choice for deciphering the parameters (desolvation energy, cation size) at the origin of the exquisite K+affinity of G-quartets^{58,59}. This wealth of information has been progressively harnessed to design ever more versatile G-based molecular tools that have found applications in the fields of soft matter, nanotechnology and chemical biology, which will be discussed in the following sections.

Nanotechnological applications

As described above, the first application developed with guanines relied on their capability to firmly chelate cations. Lipophilic SQ were thus exploited as ionophores, capable of extracting cations from water into organic solvents (most often chlorinated solvents, such as chloroform or dichloromethane). Model studies were performed with K⁺ and Na⁺, but the demonstration that G-quartets efficiently chelate radioactive metals such as caesium (137Cs+) and radium (226Ra2+) generated a tremendous strategic interest. Notably, it was shown that the isoguanosine derivative (1) could self-assemble to form nanoscale decamers that were capable of capturing $Cs^+([(1)_5Cs^+(1)_5])$ and $Ra^{2+}([(1)_5Ra^{2+}(1)_5])$, even in the presence of a million-fold excess of other cations (Na⁺, K⁺, Mg²⁺)^{21,22}, and, thus, suggesting a possible new application in the remediation of nuclear wastes.



Fig. 4 | **Structural studies of water-soluble G-assemblies. a** | Monomeric guanosine monophosphate (GMP) can self-assemble in discrete G-quartets in the presence of Na⁺ and K⁺, but the latter can also trigger the formation of micrometre-long, helical rod-like structures comprising continuous, open-ended G-quartet³⁶. **b** | The dimeric GMP c-di-GMP can form G-quadruplexes comprising four c-di-GMP molecules upon interaction

with aromatic inducers (here, thiazole orange) that intercalate between two layers of G-quartets⁴⁰. **c** | Tetrameric GMP can be assembled intramolecularly to provide water-soluble template-assembled synthetic G-quartets (TASQ) that fold into intramolecular SQ without cations and serve as a platform for investigating the G-quartet interactions of G4-ligands (here, BRACO-19)⁴¹.



Fig. 5 | **Applications of G-quartet-based devices as ionophores and ion channels. a** | Artificial ion pair (NaCl) receptor that results from the dimerization of 1,3-*alt*-calix[4]arene-templated guanosine (G)-tetramer ($\mathbf{8}$)⁶⁰. **b** | A covalently crosslinked quadruplex, comprising 16 units of the G-monomer ($\mathbf{9}$)⁶² linked via olefin metathesis, acts as a Na⁺ transporter through inclusion in a liposome membrane. **c** | The lipophilic, bis-lithocholamide-based G-dimer ($\mathbf{10}$)⁶⁵ self-associates to form synthetic G-quartets (SQ)-based nanopores that allow for K⁺ transportation through phospholipid bilayers.

Davis and colleagues later developed ionophores with improved chelating properties, providing binding sites for both cations and anions⁶⁰. The dimeric assembly of 1,3-*alt*-calix[4]arene-templated G-tetramer (**8**) indeed traps Na⁺ within the intermolecular SQ plane, while Cl⁻ is chelated by two neighbouring amide groups (FIG. 5a). This study, performed in water-saturated chloroform, highlighted the key role of water in the overall stability of the supramolecular edifice, lowering the energy required for extracting the ions from bulk water to the ionophore's binding site, participating in salt binding between the quartet layers and filling the coordination spheres of bound ions.

Lipophilic SQ were also exploited as ion channels⁶¹ — their lipophilicity means that they are readily incorporated

into membranes. Davis and colleagues constructed transmembrane pores based on a quadruplex architecture. These pores were formed by the assembly of the G-monomer 5'-(3,5-bis(allyloxy)benzoyl)-2', 3'-isopropylidene guanosine (9), followed by crosslinking of neighbouring guanines within the quadruplex architecture using ruthenium-catalysed, ring-closing metathesis⁶² (FIG. 5b). This approach leads to a covalently crosslinked quadruplex that avoids any potential problems arising from the dynamic nature of quadruplex formation/disassembly. The capacity of the resulting quadruplex, comprising sixteen units of (9), to transport Na⁺ across a liposome membrane was qualitatively demonstrated by both base-pulse assay and ²³Na NMR spectroscopic experiments. The approach of covalently capturing the formed quadruplexes was subsequently extended using a photocycloaddition strategy. Combining 2',3'-isopropylidene guanosine moieties with meta-methoxycinnamic acid provided G-monomeric units primed for crosslinking. Once assembled in SQ in the presence of K⁺, an UV-promoted [2+2] cycloaddition of the cinnamate chains led to the formation of cyclobutane covalent crosslinks with high chemical yields (>90%)63. However, the ion-transportation properties of these new suprastructures were not assessed. Matile and colleagues exploited another type of lipophilic SQ to assemble ion channels, using π -stacked supramolecular rosettes made of G-mimicking folate quartets⁶⁴. The capacity of the obtained channels to transport K⁺ across lipid bilayers highlights the importance of the dynamic nature of quartet-based channels, which uniquely allows for sterically demanding cations to be transported through topologically breathing channels, an effect termed dynamic ion tunnelling. Davis and colleagues also constructed ion channels whose structure relies on the assembly of preformed G-dimers covalently linked by bis-lithocholamide linkers (10)⁶⁵ (FIG. 5c). The capacity of these pores to enable K⁺ transport across phospholipid bilayers was quantified by conductance experiments. It was, thus, shown that the resulting SQ channels formed in this way were heterogeneous in size, with small, 2.6-Å-diameter pores (with a corresponding conductance in the pS range) that correspond to discrete SQ and large, 12-Å-diameter pores (with conductance in the nS range) that correspond to polymeric SQ (as shown in FIG. 5c). Dash and colleagues subsequently combined these different approaches, linking lipophilic G-dimers using various bridge structures (polyethylene glycol (PEG), dyes, aromatic linkers)^{66,67} to assemble channels that could embedded within phospholipid bilayers and capable of transporting cations such as K⁺ for long durations (up to 10h).

Moving beyond ion transportation, G-quartet structures have also been exploited for the construction of functional nanostructures. For instance, Dash and colleagues exposed GMP to microwave-assisted thermal treatment in order to assemble carbon dots68. While GMP itself does not possess interesting spectroscopic properties, these so-called G-dots were found to have interesting fluorescence upconversion properties and can be readily converted in fluorescent G-dot hydrogels in the presence of K⁺, which were shown to have high thermal stability (gel-sol transition temperature >70 °C) and elasticity. Meijer and colleagues used LipoGs to assemble fluorescent organic nanoparticles (ONP) (FIG. 6a). Cation-templated self-assembly of oligo(p-phenylene vinylene) (OPV) guanosines (11) leads to disc-shaped octameric nanostructures (1.5 nm high, 8.5 nm wide), with improved fluorescence emission properties in comparison with isolated guanosines (with a twofold to threefold enhancement of quantum yield in organic solvents) and thermal stability (melting temperature >70 °C)⁴⁵. Mirkin and colleague also exploited G-quartet formation to assemble metallic nanoparticles (FIG. 6b). A disperse solution of 10-nm gold nanoparticles coated with 3'-thiopropyl-GMP (12) were combined with

a series of short, G-rich oligonucleotides to produce, upon Na⁺ addition, micrometre-long architectures that displayed increased plasmon coupling as a result of reduced interparticle distance and higher thermal stability (aggregate-sol transition temperature >60 °C)⁶⁹. Wu and colleagues assembled even larger nano-objects by subjecting concentrated GMP solution (33 wt%, i.e. 800 mM) to weakly basic conditions (pH = 8), which results in the formation of Na⁺-containing nanocylinders (26 Å diameter, up to 30 nm long)⁷⁰. Qu and colleagues later demonstrated that replacing Na⁺ for Sr²⁺ enabled the production of micrometre-long nanowires from diluted GMP solution (20 mM), in a reversible, controllable manner (nanowires were broken upon the addition of EDTA)71. Decorating these fibres with three dyes, i.e. thioflavin T, thiazole orange and pyronin Y, that constitute a fluorescence resonance energy transfer (FRET) system, the same authors reported on the use of GMP-based nanowires as light-harvesting material (FIG. 6c). Once assembled in the nanostructure, light $(\lambda_{ex} = 450 \text{ nm})$ is absorbed by thioflavin T, transmitted by FRET first to thiazole orange and then pyronin Y $(\lambda_{em} = 573 \text{ nm})$, generating a photocurrent that demonstrates the functional relevance of triple-dye-doped nanofibres as light-harvesting antennae⁷². The electronic properties of G-based suprastructures were further studied by Wasielewski and colleagues, who investigated the properties of guanines with electron-acceptor moieties such as perylene-3,4,9,10-bis(dicarboximide) (PDI) derivatives. The resulting GPDI self-assembles into octamers upon interaction with K+ (FIG. 6a). Femtosecond transient absorption spectroscopy was used to demonstrate that [(GPDI)K⁺(GPDI)] undergoes efficient charge separation upon photoexcitation at $\lambda_{ex} = 525$ nm. The lifetime of the charge-separated intermediate in the octameric assembly was determined to be 1.2 ns, as compared with 98 ps for the monomeric GPDI73. Using a dimeric system with two guanosines appended to the core perylene diimide, the same authors constructed SQ organic frameworks and showed their ultrafast production of long-lived mobile charges (radical ion pairs, or polarons) upon photoexcitation ($\lambda_{ex} = 500 \text{ nm}$; FIG. 6d). The excellent electrical conductivity of these frameworks was exploited to assemble a SQ-based Li-ion battery, which proved to be robust (<7% loss of efficiency after 300 charge/discharge cycles)74.

G-quartets also served as a platform for interacting with small-molecule partners to endow the resulting G-quartet/ligand complex with specific properties. Two examples were discussed above: the interaction of thiazole orange with a c-di-GMP octamer (FIG. 4b), in which the resulting quadruplex/ligand complex is fluorescent (and, thus, detectable), and the interaction of TASQ with G4-ligands (FIG. 4c), which allows for quantifying in vitro the parameters that govern the ligand/ target interaction. Combining these two approaches, Sintim and colleagues developed a bioinspired catalytic system for the oxidation of 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS; FIG. 7). In the presence of an intercalator (proflavine), c-di-GMP forms a quadruplex that externally binds hemin (protoporphyrin IX iron(III) chloride)75, a well-known cofactor of

Fluorescence upconversion

this fluorescence spectroscopy (also known as sum frequency generation), is a nonlinear, high-resolution technique characterized by an emission wavelength that is shorter than the excitation wavelength.

Organic frameworks

three-dimensional organic structures in which building blocks are associated either through covalent bonds (covalent organic framework, or COF) or metal-mediated coordination (metal organic framework, or MOF).



Fig. 6 | **The nanotechnological applications of synthetic G-quartets-based nanostructures. a** | Decorated lipophilic guanosines self-assemble into octamers with enhanced properties, i.e. enhanced fluorescence for (**11**)⁴⁵ and charge separation under photoexcitation for guanine perylene-3,4,9,10-bis(dicarboximide) (GPDI)⁷³. **b** | Gold nanoparticle assembly triggers by (**12**)⁶⁹, which provides micrometre-long architectures with increased plasmon coupling. **c** | Guanosine monophosphate (GMP)-based nanowires that act as light-harvesting antenna upon association with fluorescence resonance energy transfer (FRET) dyes triplets⁷², i.e. thioflavin T (ThT), thiazole orange (TO) and pyronin Y (PY). **d** | G-quadruplex organic frameworks that produce long-lived mobile charges upon photoexcitation, making them prototypes of a G-quartet-based Li-ion battery⁷⁴. OPV, oligo(*p*-phenylene vinylene); PDI, perylene-3,4,9,10-bis (dicarboximide); SQ, synthetic G-quartets.

hemoproteins, such as horseradish peroxidase (HRP), and a key biochemical tool in the development of the enzyme-linked immunosorbent assay (ELISA). The catalytic activity of G-quartet-containing biomolecules, i.e. quadruplex-DNA (G4-DNAzyme), was discovered during investigations that aimed to identify aptamers for catalysing porphyrin metalation. Aptamers with a high affinity for N-methyl mesoporphyrin IX (NMM) were investigated, given that NMM is a transition-state analogue for porphyrin metalation and a potent inhibitor of ferrochelatase, an enzyme that catalyses the insertion of iron into protoporphyrin IX (the last step of haem biosynthesis). Through systematic evolution of ligands by exponential enrichment (SELEX) experiments, Sen and colleague identified a short, 24-nt DNA referred to as PS5.M that strongly binds to NMM and catalyses the metalation of mesoporphyrin IX76,77. Importantly, it was shown that the catalytic activity of PS5.M is inhibited by hemin through the formation of a high-affinity PS5.M/hemin complex. This observation drove the authors to assess whether it could elicit hemoprotein-like

properties, notably peroxidases. Upon addition of a stoichiometric oxidant (hydrogen peroxide, H₂O₂), the complex was found to accelerate 250-fold the oxidation of the chromogenic substrate ABTS78. In light of the G-richness of PS5.M (d[5'GTG₃TCAT₂GTG₃TG₃ TGTG₂^{3'}]) and the K⁺-dependency of its catalytic activity, the authors wisely mused about its quadruplex structure79, which offers a defined binding site to hemin that becomes catalytically competent once nestled within this site only. It is intriguing that, more than three decades after its discovery, the precise mechanism of quadruplex DNAzyme remains unclear⁸⁰⁻⁸², despite hundreds of new applications reported each year (for relevant reviews, see REFS⁸³⁻⁸⁵). Inspired by Sintim's work, Spinelli and colleagues further exploited the intramolecular assembly of TASQ to provide versatile catalytic systems (TASQzyme) that could be applied with a variety of pre-catalysts, cofactors and substrates⁸⁶. Monchaud and colleagues used the non-anionic TASQ (e.g. DOTASQ; FIG. 7) (in place of anionic DNA or c-di-GMP) to replace the poorly soluble hemin cofactor with the more practically



Fig. 7 | **DNAzyme-type catalysis performed by G-quartet molecular tools.** Catalytic guanosine (G)-quartets comprising either proflavine-promoted c-di-GMP octamer⁷⁵ or the intramolecularly folded DOTASQ⁸⁷ upon interaction with hemin, which makes the resulting G-quartet/hemin complex catalytically competent to oxidize the chromogenic 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) in the presence of H_2O_2 (as the stoichiometric oxidant), in a DNAzyme-type manner⁸⁵. SQ, synthetic G-quartets.

Logic gates

in electronics, a logic gate is a device that performs a logical operation on one or two inputs and produces a single output. Logic gates are primarily implemented as switches of different possible natures (AND, OR, NOT, INHIBIT etc.) in digital circuits. convenient 5,10,15,20-tetrakis(4-sulfonatophenyl) porphyrinato iron (III) chloride (FeTPPS)87. The construction of catalytic G-quartets is, however, not limited to assembly from G-dimers and G-tetramers: recently, Davis and colleague showed that GMP itself could assemble to form an efficient thermosensitive catalyst for DNAzyme-like reactions, offering higher catalytic rates at low temperature, owing to an improved stability of GMP-based G-quadruplex structure at 10 °C versus 30 °C (REF.88). Dash and colleagues reported on the design of catalytic G-quartet-based hydrogels formed by boronate-ester-functionalized guanosines that act as logic gates. K⁺ and Pb²⁺ are the on and off inputs, respectively, and the product of the oxidation of 3,3',5,5'-tetramethylbenzidine is the output, thanks to its characteristic absorbance signal at 652 nm that provides a convenient readout of the efficiency of the operation⁸⁹. Pei and colleagues exploited a similar approach to design bioelectronics nanodevices (i.e. an electrochemical glucose sensor based on the concerted action of glucose oxidase and hemin-mediated catalytic G-quartets embedded in a hydrogel matrix)90.

G-quartet-based soft matter

Soft matter refers to materials whose properties occupy the tenuous frontier between the solid and liquid states. The study of soft matter was popularized by de Gennes, whose work was recognized by the award of the Nobel Prize in Physics 1991 (REF.⁹¹). These materials have low elastic moduli, enabling them to withstand thermal fluctuations and external deformations, thanks to internal structural rearrangements that allow for energy dissipation in response to external stress⁹². They can be produced from a variety of organic and inorganic molecular entities (e.g. clay, metal oxides, synthetic and natural polymers and proteins) and encompass a wide diversity of systems, including colloid suspensions, emulsions, foams, surfactants, liquid crystals or gels.

The pioneering works on the ability of GMP to form hydrogels^{7,8,10} cast a bright light on G-quartets as soft-matter building blocks⁹³. For example, Gottarelli et al. studied the formation of columnar aggregates by the lipophilic 3',5'-didecanoyl-2'-deoxyguanosine (**2**)^{94,95} and sought to characterize them both structurally and stereochemically. They noticed that



Fig. 8 | **Soft matter based on G-quartet molecular devices. a** | Amphiphilic guanosine (G; **2**) that self-assembles into a lyotropic liquid crystal triggered by a K⁺-promoted synthetic G-quartets (SQ) formation⁹⁶. **b** | Cation-free SQ formed by adamantyl-conjugated guanines (**13**)⁹⁹, whose SQ/ribbon topological transition is controlled by the concentration. **c** | Dynamic hydrogel (or dynamers) formed by hydrazide-based guanines (**14**)¹⁰⁰ and controlled by cryptand [2.2.2] and sequential acid/base addition. **d** | Water-assisted polymerization of siloxane-conjugated guanines (**15**)¹⁰³ to form covalent crystalline nanorods.

Lyotropic liquid crystal

property of an amphiphilic material (mesogen) that exhibits phase transitions upon dissolution in a suited solvent, in a concentration-dependent manner. lyotropic liquid crystal phases were formed in heptane in the presence of potassium picrates⁹⁶. At high dilution (<6% w/w), an isotropic phase was observed, while medium (6–16% w/w) and high (40–50% w/w) concentrations led to a left-handed cholesteric phase and a hexagonal phase, respectively (FIG. 8a). X-ray measurements confirmed the columnar nature of the G-quartet-based self-assemblies with characteristic peaks at 3.4 Å corresponding to the stacking of G-quartets and an additional calculated radius of 12 Å for the hexagonal phase, fitting with the G-quartet radius. However, in ion-free conditions, **2** and other LipoG derivatives also exhibit lyotropic properties in both toluene and chloroform, in this case due to the formation of G-ribbons⁹⁷. Recently, Kato and colleagues reported on thermotropic liquid-crystal formation with C8-oligothiophene guanine derivatives. In this case, both the nature of the phases and the phase transitions were modulated by potassium ion concentration. As well as the hexagonal columnar phases based on G-quartet assemblies (as already reported for **2**),

Storage moduli

indications of the ability of the gel to withstand and store deformation energy in an elastic manner, being dependent on its crosslinking state.

Thixotropy

property of a material (gel, fluid), which is thick or viscous under static conditions, to become thin and liquid when shaken or sheared. G-ribbon formation leads to bicontinuous cubic phase rarely observed for π -conjugated liquid crystals⁹⁸. G-quartets have also been used to formulate organogels and hydrogels, assembled in organic solvents or aqueous media, respectively. As an example, Yi and colleagues demonstrated that 5'-(adamantylcarbamoyl)propionate-2', 3'-isopropylidene-guanosine (13) can form an organogel even in the absence of cations when working above the critical gelation concentration (>50 mM). The sol-gel transition results from a structural change from G-quartets (assembled at low concentration, <10 mM) to type A G-ribbons (FIG. 8b). Gelation occurred when ribbons self-assembled into micrometre-width hexagonal columnar structures; a morphological change can be triggered by ultrasonication, which favours twisted-flower-like structures99. Liu and colleagues observed a similar behaviour with a guanine functionalized with a 3,4,5-tris(n-dodecan-1-yloxy)benzohydrazide (14): the quartet-to-ribbon structural transition was solvent-dependent, occurring above 17% w/w in toluene and above 29% w/w in chloroform. This transition was harnessed to build a dynamic organogel. In the presence of cryptand [2.2.2], which is known to chelate K⁺ ions, sequential additions of trifluoroacetic acid and triethylamine trigger switching between the G-quartet formation (sol state) in the presence of base and the G-ribbon formation (gel state) in the presence of acid¹⁰⁰ (FIG. 8c). The strategy of using cryptands to modulate G-ribbon/G-quartet interconversion was previously described in organic solution by Spada and colleagues with LipoG (2)101 and a 5'-O-oligothiophene guanine derivative¹⁰². Higher-order, self-organized, hybrid organic/inorganic materials have also been built on G-quartet-based organogels formulated from a guanine-siloxane derivative (15) in the presence of K⁺ or barium (Ba²⁺) ions¹⁰³. The polymerization of – SiR₂-OEt into -SiR₂-O-SiR₂- was triggered by water, leading to higher-ordered hybrid materials with micrometric crystalline-rod morphologies (FIG. 8d). Similarly, self-organized organic polymers that exhibit microfibre or microspring morphologies can be accessed from a 5'-O-methacrylate guanosine derivative, with the morphology switching dependent on cationdriven G-ribbon or G-quadruplex adopted assembly, respectively104.

The majority of G-quartet-based soft matter described to date focuses on the formation of hydrogels. From the initial pioneering observations of the GMP gelation process^{7,8,10}, a vast amount of research has been conducted in order to decipher the parameters that govern the formation of hydrogels and improve the scope of their applications. GMP-based gels have mostly been formulated with monovalent or divalent cations (see above); however, the use of trivalent lanthanide ions, including lanthanum (La³⁺), cerium (Ce³⁺), europium (Eu³⁺) and terbium (Tb³⁺), led to SQ-based nanofibrillar networks in acidic conditions¹⁰⁵. As a result of its high aqueous solubility, GMP alone does not form a gel at neutral pH. However, upon addition of phosphate-free guanosines, gelation occurs and the modulation of the GMP/guanosine ratio can be used to fine-tune the resulting material properties^{106,107}.

These gels exhibit unusual thermoassociative behaviour, with the sol-gel transition taking place above a specific temperature¹⁰⁸. In the presence of different monovalent cations (from Li⁺ to Cs⁺), guanosine alone can also form a hydrogel, but it is stable for only a few minutes or hours. This instability can be addressed by the addition of 0.7-1.5 molar equivalents of lipophilic 2',3',5'-tri-O-acetylguanine (16), which enhances the gel lifetime from hours to years and also increases the sol-gel transition temperature (from 50 °C to >60 °C)¹⁰⁹ (FIG. 9a). Another possibility is to formulate the gel with metal ions such as silver (Ag⁺), which increases the stability (from hours to months) and the storage moduli (up to 8 kPa) and endows the resulting gel with thixotropy¹¹⁰. Addition of isoguanosine is another alternative to turn guanosine gels into thixotropic co-gels exhibiting very short recovery times (<20 s)¹¹¹.

Davis and colleagues thoroughly exploited guanosine-borate assemblies¹¹², which showed high propensity to form stable hydrogels (with storage moduli >10 kPa at 0.2 wt%; FIG. 9b). These gels have been used to incorporate a wide variety of chemicals, notably fluorescent dyes. Even at extremely low concentration $(2.5 \times 10^{-3} \text{ molar})$ equivalents), these probes impact the hydrogel properties - decreasing gel stiffness in the case of, e.g. rhodamine B and Rose Bengal, or increasing it in the presence of, e.g. thioflavin T or thiazole orange¹¹³. Guanosine-borate systems were also used in the preparation of photoreversible hydrogels, in which the photoswitch E-4,4'dioxyethylpiperidinium iodide-azobenzene controls the G-quartet-based assembly ($\lambda_{ex} = 436 \text{ nm}$) and disassembly $(\lambda_{ex} = 365 \text{ nm})^{114}$. Lehn and colleague developed smart, G-quartet-based hydrogels, using water-soluble G-dimer (17). As above, the sol-gel equilibrium of these K⁺-promoted dynamic polymers can be fully controlled by sequential addition of cryptand [2.2.2] and HCl to control the availability of K+ to promote G-quartet formation¹¹⁵. Another dynamer was developed to exploit the reactivity of monomeric guanosine-5'-hydrazide (18) with a library of aldehydes, in order to assemble acylhydrazone-based hydrogels capable of dynamically self-selecting the most stable gel under the experimental conditions¹¹⁶⁻¹¹⁸ (FIG. 9c). Barboiu and colleagues reported a double-dynamer system based on the reversible reaction between guanosines and a bisiminoboronic acid linker to form bisiminoboronate-G-dimers, whose equilibrium between the cation-free ribbon state and K⁺-promoted G-quartet state led to the production of two types of functional membrane films¹¹⁹.

Interesting biotechnological and biological applications have been reported with G-quartet-based hydrogels. Dash et al. used Ag⁺ to formulate hydrogels that enable the incorporation of cytochrome *c* into GMP-based gels¹²⁰. Zhao and colleagues described the preparation of an antimicrobial hydrogel (active against Gram-negative bacteria) by incorporating 2'-deoxy-2'fluoroguanosine (**19**)¹²¹ (FIG. 9a). The controlled release of bioactive compounds from G-quartet-based hydrogels has been described as a method for drug delivery. For example, the release of vitamin C and the G-containing drugs acyclovir and ganciclovir has been achieved using hydrogels based on guanosine-5'-hydrazide¹²²,



Fig. 9 | **Supramolecular hydrogels based on G-quartet assemblies. a** | Guanine derivatives (**16**)¹⁰⁹, (**17**)¹¹⁵, (**19**)¹²¹, (**20**)¹²³ and (**24**)¹²⁶ used as building blocks for hydrogels. **b** | Guanosine–borate-based hydrogels used for incorporating a wide variety of chemicals¹¹². **c** | Dynamic self-selection of aldehydes from (**18**) that provides the most stable gel¹¹⁸. **d** | Following gel formation, the cyclization of (**21**) to (**22**) triggers gel weakening and collapse, allowing the controlled release of acyclovir incorporated into the stacked guanosine (G)-quartet assemblies¹²⁴. **e** | Biological application of a platinum–G-quartet-based hydrogel exhibiting photoactivatable anticancer properties¹²⁵.

guanosine/bromoguanosine (20) (1/1) mixture¹²³ or 5'-deoxy-5'-iodoguanosine $(21)^{124}$. In the latter case, an intramolecular cyclization that forms 5'-deoxy-N3,5'-cycloguanosine (22) leads to gel breakdown, triggering the release of acyclovir (>80% release after 3 days) that had previously been incorporated into the stacked G-quartet assemblies through its guanine moiety (FIG. 9d). Sadler and colleagues further exploited this approach, incorporating a guanosine-borate Pt(IV) complex (23) into a guanosine-borate hydrogel to exploit the photo--activatable anticancer properties of platinum¹²⁵. Under visible-light irradiation ($\lambda_{ex} = 465 \text{ nm}$), the photoreduction of Pt(IV) to cytotoxic Pt(II) combined with the release of azidyl radicals (FIG. 9e) triggers higher cytotoxicity in cisplatin-resistant A2780cis human ovarian cancer cells (IC₅₀ = $3 \mu M$) than cisplatin and chlorpromazine (IC₅₀ > 100 μ M and 6 μ M, respectively), with a better selectivity towards normal MRC-5 human fibroblast cells (IC₅₀ > 50 μ M).

Rowan and colleagues described the application of hydrogels derived from 8-methoxy-2',3',5'-tri-O-acetylguanosine (24) as a scaffold for tissue engineering. C166 endothelial cells were incubated in cell culture medium DMEM/24 mixtures (between 0 wt% and 2 wt%), the addition of gelatin (1 wt%) providing a tridimensional co-gel within which cells can be incorporated, as observed by confocal microscopy¹²⁶. The choice of benzene-1,4-boronic acid instead of simple boronic acid to form a guanosine-borate hydrogel provides an efficient support for human dermal fibroblast growth with cell viability >73% when formulated with both K⁺ (triggering the G-quartet formation) and magnesium (crosslinking the anionic borate moieties ions)¹²⁷. Altogether, these studies confirm the versatility of G-quartet-based soft matter and highlight their high potential as biomaterials.

Applications in chemical biology

As it may have become apparent from the discussion above, increased understanding and popularity of G-quartets as supramolecular building blocks has also increased our understanding of these structures in their natural setting - within DNA and RNA quadruplexes. The prevalence of G4s in the human genome, transcriptome and ncRNAome (G4ome) is now firmly demonstrated thanks to the recent and spectacular development of sequencing technologies¹²⁸ (FIG. 1): G4-seq identified >716,000 potential DNA quadruplex-forming sequences (QFS) in the human genome¹²⁹ and G4 ChIP-seq decreased this number to 10,000 G4 sites, because of the topological restriction imposed by the chromatin environment¹³⁰; rG4-seq identified >13,000 QFS in the human transcriptome (polyA-enriched RNA)¹³¹, while G4RP-seq extended the analysis to the whole transcriptome and ncRNAome, demonstrating that G4s are pervasive structures in functional human cells¹³². Therefore, the prevalence of G4s and their established functional relevance16,17 have made them new stars in genetics and promising targets for therapeutic interventions. Specially dedicated G4-ligands¹³³⁻¹³⁵, such as quarfloxin¹³⁶, BRACO-19 (REF.¹³⁷), PhenDC¹³⁸ and pyridostatin¹³⁹, were developed in the hope of gaining control over the many cellular processes in which G4s are involved. Most, if not all, programmes that have attempted to identify promising new G4-ligands have chemical scaffolds primarily developed as duplex DNA intercalators (acridine, quinoline, etc.). This approach is somewhat surprising, given that binding to duplex DNA is the major unwanted interaction that would need to be avoided for a G4-ligand to be considered promising. That said, successful strategies have been devised that avoid duplex interaction, such as using ligands with large to very large aromatic cores to maximize overlap with the external G-quartet of a G4 structure and avoid duplex intercalation. Another approach that has emerged — as a result of the many structural studies on quadruplexes - is the use of synthetic G-quartets as bioinspired G4-ligands. This approach builds on the observation that the stability of quadruplex structures is frequently related to the number of constitutive G-quartets, and there is a like-likes-like recognition between natural and synthetic G-quartets (FIG. 10).

In order to circumvent the challenging kinetics associated with the assembly of four isolated guanines, the first biomimetic ligands developed were tetrameric TASQ, such as the neutral DOTASQ^{140,141} (FIG. 7). Further work led to the development of the first smart quadruplex ligands, which assemble only in the presence of their native G-quadruplex targets, for example, the cationic PNADOTASQ142,143 (FIG. 10a). PyroTASQ has been described as a 'twice-as-smart' quadruplex ligand^{144,145} (FIG. 10a), referring to the fact that it is both a smart quadruplex ligand and a smart, turn-on fluorescent probe. The exquisite selectivity of these ligands for their targets is a result of a unique G4-triggered intramolecular rearrangement of the ligand. TASQ ligands have been instrumental in the detection and identification of DNA/RNA quadruplexes in human cells using optical imaging (for example, N-TASQ was used as a probe to directly visualize DNA/RNA quadruplexes in human cells; FIG. 10a,b)^{146,147} and sequencing studies (BioTASQ was used for the pull down of RNA quadruplexes containing sequences from living human cells, enabling their identification through to the G4RP-seq protocol (G4-RNA-specific precipitation followed by sequencing); FIG. 10a,c)^{132,148}. Together, these studies have both confirmed the existence of quadruplexes in functional human cells and provided tools to assess the ligand-induced changes in the quadruplex landscape upon therapeutic treatments.

Intermolecular SQ have also been used as biomimetic ligands. Rivera and colleagues reported on the use of quartets self-assembled from the 8-aryl-2'deoxyguanosine derivative (**25**) and showed that this cationic guanine spontaneously folds into K⁺-promoted or Na⁺-promoted hexadecamers, which are subsequently sandwiched between two quadruplex units $[G4((25)_4K^+(25)_4K^+(25)_4)G4]^{149}$ (FIG. 10d). These unusual, nanometre-long structures, whose stability is a result of extended networks of π -stacking and Coulombic interactions, describe a new possible G4-ligand/quadruplex binding mode and make **25** SQ the first example of a biomimetic, self-assembled ligand. Dash and colleagues also reported on the use of G-monomers and G-dimers



Fig. 10 | **The chemical biology applications of SQ. a** | Chemical structure of several template-assembled synthetic guanosine (G)-quartets (TASQ) used as biomimetic G-quadruplex ligands, i.e. ^{PNA}DOTASQ¹⁴² and PyroTASQ¹⁴⁴, N-TASQ^{146,147} and BioTASQ^{132,148}. **b** | Schematic representation of the detection of quadruplexes in human cells using the twice-as-smart N-TASQ ligand. **c** | Schematic representation of the use of BioTASQ as molecular bait to fish quadruplexes out of human cells prior to being identified by sequencing (G4RP protocol). ** and *** represent the statistical analysis (T test) of the results, *p*-values (with ** *p* < 0.01 and *** *p* < 0.001)¹³². **d** | Self-assembled G4-ligand resulting from the multimerization of (**25**)¹⁴⁹ that is subsequently sandwiched between two quadruplex units.

(dansyl monoguanosine and dansyl diguanosine) as G-quadruplex-interacting agents¹⁵⁰. It is thought that these agents are groove binders (rather than themselves forming a quartet structure), but they were found to interact efficiently with the quadruplex found in the promoter region of the Myc oncogene (a gold-standard target for anticancer strategies) and capable of downregulating its expression in cancer cells.

Conclusions

Guanine was, is and will remain a fascinating study subject and likely has many functionalities that are yet to be discovered. Described above are just some of the applications that have been developed in the past two decades, and highlight an apparent disconnect between the simplicity of the guanine chemical scaffold and the diversity of fields in which guanines and their derivatives provide practical solutions. The careful selection of derivatives (guanine itself, guanosine, deoxyguanosine, GMP, etc.), combined with suitable chemical modifications, has enabled the assembly of a toolbox that appears to be virtually unlimited. Chemical modifications mean that guanine is useful in a variety of experimental conditions (organic or aqueous environments) and responsive to external stimuli (cation, pH, light and temperature), which translates into programmable supramolecular properties. However, it is also immediately apparent that we are only beginning to assess the functional diversity of guanine in its entirety and to harness its supramolecular properties in a fully controllable manner.

To move up a gear, we believe that this field will have to shortly meet the following milestones.

Better predictions. Efforts must be invested to design G-based systems that will strictly obey predefined rules in terms of topological pluralism and stability. We need to understand and harness these rules with the help of in silico models to predict and encode the properties of G-based devices in the very structure of their basic guanine units.

Improved chemical strategies. The commercial accessibility of guanine derivatives sustains the use of G-based molecular tools in various research fields. However, the

many examples discussed above highlight that only exquisitely tailored guanines provide devices with controllable properties. The relevance of a G-based molecular toolbox is, to date, far too dependent on high-tech chemistry. This might prevent academia-to-industry transitions in the short term.

Alternative chemical approaches. The supramolecular nature of this field of research must be further exploited, notably through alternative chemical strategies, such as dynamic combinatorial chemistry. This approach will undoubtedly offer alternative solutions to technical conundrums and improve the modularity and diversity of G-based devices.

New fields of applications. Beyond the aforementioned examples, more applications might benefit from G-based materials, from personalized medicine (for example, the use of G-rich aptamers as therapeutics) to climate-change issues (for example, G-polymers harnessing solar energy). This will require the input of scientists from diverse research domains, making this field a resolutely multidisciplinary research area.

Validated biotech prototypes. Important challenges will have to be met in terms of scalability and cost to translate this research into the industrial arena. The advent of successful applications in the next years will help take the last few steps towards long-awaited demonstration of the industrial significance of G-based molecular devices.

Hammarsten initially described guanine as a "curious nucleic acid"⁶; we would wager that he was very far from thinking that, 125 years later, scientific communities from different origins (supramolecular chemistry, polymer, soft matter, bioelectronics, catalysis and chemical biology) would become curious about this nucleic acid. One thing is certain: no one is more adventurous than the scientist who tries to follow guanine on its way across various chemical lands, along a route whose trajectory is as yet unknown. It is a fascinating, transdisciplinary chemical adventure in a yet undiscovered country from which no wanderer returns.

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Author contributions

All authors contributed equally to the preparation of this manuscript.

Competing interests

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