Prokaryotic Argonaute Proteins as a Tool for Biotechnology

E. V. Kropocheva*, L. A. Lisitskaya*, A. A. Agapov*, A. A. Musabirov*, A. V. Kulbachinskiy*, and D. M. Esyunina*, *

*Institute of Molecular Genetics, National Research Centre “Kurchatov Institute,” Moscow, 123182 Russia
*e-mail: es_dar@inbox.ru
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Abstract—Programmable nucleases are the most important tool for manipulating the genes and genomes of both prokaryotes and eukaryotes. Since the end of the 20th century, many approaches were developed for specific modification of the genome. The review briefly considers the advantages and disadvantages of the main genetic editors known to date. The main attention is paid to programmable nucleases from the family of prokaryotic Argonaute proteins. Argonaute proteins can recognize and cleave DNA sequences using small complementary guide molecules and play an important role in protecting prokaryotic cells from invading DNA. Argonaute proteins have already found applications in biotechnology for targeted cleavage and detection of nucleic acids and can potentially be used for genome editing.

Keywords: programmable nucleases, Argonaute proteins, guide DNA, biosensors, genome editing
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MAIN GROUPS OF PROGRAMMABLE NUCLEASES

Targeted changes in the DNA sequence at the level of specific genes and regulatory regions of the genome are highly important for fundamental studies of the mechanisms of genome functioning and the role of individual proteins, as well as for solving applied problems, from the obtaining of producer strains to the treatment of genetic diseases. The most important tool in these studies are programmable nucleases, enzymes that can be directed to a specific region in the genome to make changes in the DNA sequence. Catalytically inactive variants of programmable nucleases can be used to edit epigenetic marks and regulate gene expression by attracting various functional domains to certain regions in the genome and to visualize the target regions in cells. Programmable nucleases are also used to identify target sequences in biological samples and to cleave nucleic acids in vitro.

To date, several groups of programmable nucleases of natural and synthetic origin are known, differing in the mechanism of DNA recognition and cleavage and the potential for practical application. In all cases, the introduction of breaks in the target DNA region can then lead to the loss of the functionality of this gene, or to the required change in its sequence during repair. Two large groups of such enzymes can be distinguished: nucleases that recognize the cleavage site using DNA-protein interactions, and nucleases that recognize DNA through an associated short complementary RNA or DNA. Although historically the enzymes of the first group were found and used earlier, the enzymes of the second group have much greater flexibility in recognizing target DNA regions and have been used as the main tools for genome editing during last years. A comparison of different classes of programmable nucleases is shown in Fig. 1.

Meganucleases

Meganucleases recognize long (from 12 to 44 bp) DNA regions [4, 5]. Naturally occurring meganucleases belong to the group of homing endonucleases that promote mobility of transposable elements. Meganucleases cleave DNA strands in the area of four central nucleotides in the recognition site with the formation of a double-strand break with 3’-terminal 4-nucleotide overhang [6, 7]. The small size of these proteins with a long DNA recognition site has drawn attention to it as tools for genome manipulation. The limitation is that the recognition site must be artificially introduced into the genome. By the early 2000s, several hundred endonucleases with different recognition sites had been discovered and described [5], and enzymes with altered specificity were created [8–10].

Chimeric Nucleases

The idea of creating chimeric nucleases by combining the nuclease domain of one protein and the DNA-recognition domain of another protein in one molecule was proposed in 1994 by Kim & Chandrasegaran [11]. The researchers combined the nuclease domain of the Fla-
vobacterium okeanokoites FokI restriction endonuclease, which does not have specificity for a certain DNA sequence, and the homeodomain of the Drosophila melanogaster Ultrabithorax regulatory protein, which recognizes a 9 nucleotide long DNA sequence. The authors also proposed variants of DNA-recognition domains from other proteins, which were indeed later used to create new chimeric nucleases.

**SGN**

SGN (structure-guided endonuclease) is a structure-specific endonuclease. Nucleases can be targeted to a specific location in the genome by recognizing specific structures in DNA. An example is the nuclease consisting of the flap endonuclease (flap endonuclease-1) FEN1 of the archaea Archaeoglobus fulgidus and the FokI nuclease domain [12, 13]. To introduce a break at the desired location in the DNA, DNA guides are used that are complementary to the desired site, with the exception of one 3'-terminal nucleotide. This nucleotide is recognized by FEN1 within the SGN, then FokI introduces a DNA break at a distance of 9–10 nucleotides from the guide 3' end. However, SGN can also recognize 3'-protruding single-stranded fragments that arise during repair and replication. This leads to significant non-target rearrangements of the genome [12, 13]. Along with low efficiency, this has become a significant limitation for the development of SGN technology.

**Zinc Fingers**

The zinc finger domain (ZF) was first described by J. Miller et al. in 1985 [14] while studying the transcription factor TFIIIA from frog oocytes. TFIIIA consists of 9 repeats each containing 30 amino acid residues (a.a.), including two conserved histidines and two cysteines; polar and basic amino acids are concentrated at the top of each “finger” responsible for interactions with nucleic acids [14, 15]. One zinc finger is responsible for recognition of a 3-nucleotide sequence [16, 17]. The first site-specific nuclease containing zinc fingers as a DNA recognition domain and FokI nuclease domain (ZFN, zinc finger nuclease) was obtained and described by Y. Kim et al. in 1996 [18].
The recognition of individual nucleotides by individual amino acids within the zinc finger and the ability of the ZF to work as independent modules in recognizing extended DNA sequences make it possible to use these domains to program artificial protein tools designed to work with nucleic acids. Studies of chimeric ZFN molecules in Drosophila and mammalian cells revealed their high cytotoxicity due to nonspecific DNA recognition and cutting (off-target effect) [16]. An obvious way to increase the target recognition specificity with ZFN is to increase the number of zinc fingers. However, Y. Shimizu et al. [19] found that the activity of nucleases with five or six zinc fingers is much lower compared to proteins with one such activity. The activity of nucleases with five or six zinc fingers is much lower compared to proteins with one such activity. Furthermore, ZFN specificity was increased by amino acid modifications in the amino-terminal parts of the zinc fingers of the nuclease domain. Thus, the creation of functional nucleases with multiple zinc fingers is a difficult task.

**TALEN**

TALEN (transcription activator-like effectors nuclease) is an effector nuclease similar to transcription activators. TALE is a large family of transcriptional activators that belong to the virulence factors of phytopathogenic bacteria of the genus *Xanthomonas*. In 1989, U. Bonas et al. [20] showed that these proteins contain repeating elements. Repeats have the same sequence, except for amino acids at positions 12 and 13 (repeat variable di-residues, RVDs). This pair of amino acids is responsible for the specific recognition of a single nucleotide in DNA [21–23]. The modular structure and specificity to individual nucleotides of TALE DNA-recognition domains opens up the possibility of their application as a component of chimeric nucleases. The first TALE-based chimeric nucleases (TALEN–TALE nuclease) were constructed on the basis of a repeat-containing fragment of the TALE protein and the FokI nuclease domain [24].

**Cas Nucleases**

In recent years, Cas nucleases, which are part of the CRISPR–Cas systems of prokaryotic adaptive immunity (clustered regularly interspaced short palindromic repeats), have been most widely used in genomic editing in recent years. CRISPR–Cas systems have a modular organization. The two main components are the module responsible for adaptation (inserting new spacers into the CRISPR cassette) and the effector module responsible for interference. The modules are represented by complexes of Cas proteins, the genes of which are located in the same operon. Based on the structure of the effector module, two classes of CRISPR–Cas systems are distinguished. In Class 1 systems, the effector module is represented by a multisubunit complex, which consists of several Cas proteins. In Class 2 systems, effector module function is performed by one large protein with several domains. Class 1 includes CRISPR–Cas systems of types I, III, IV, and Class 2 includes II, V and VI [25].

Class 2 effector modules are the most interesting for practical application, since they are represented by a single protein molecule with nuclease activity.

Cas9 nuclease from the bacterium *Streptococcus pyogenes* (SpCas9), a representative of type II CRISPR–Cas systems, has been most thoroughly studied. Two RNAs are required for Cas9 activity: crRNA (CRISPR RNA) containing the spacer sequence and tracrRNA (trans-activating crRNA). Cas9 loaded with crRNA–tracrRNA introduces a double-strand break in the target DNA sequence corresponding to the spacer. For practical applications of Cas9, crRNA and tracrRNA can be combined into one RNA molecule, single guide RNA (sgRNA). Cas9 is a large DNA endonuclease (SpCas9 consists of 1368 a.a.), introducing a double-strand break with the formation of a blunt end within the region recognized by sgRNA, at a distance of 3 nucleotides from its right end. DNA cleavage is carried out by two nuclease domains. The HNH-like domain cuts the DNA strand complementary to the guide RNA (target strand). The RuvC-like domain has an RNase H-like fold and cuts the opposite strand (non-target strand) [26]. In early 2013, three studies were published that showed the possibility of using Cas9 as a programmable DNase to introduce changes in the genome in human cell cultures [27–29]. This marked the beginning of the Cas9 nuclease use in science and biotechnology.

sgRNA allows directing Cas9 to any DNA sequence containing a PAM motif (protospacer adjacent motif) to the right of the target sequence. PAM recognized by SpCas9 is a trinucleotide 5'-NGG-3', where N is any of the four DNA nucleotides [30], the length and sequence of this motif may vary for different proteins of the Cas family. To expand the range of sequences available for Cas9 recognition, variants of the enzyme with altered PAM specificity were constructed [31, 32], and Cas9 proteins from other bacteria were also used [33–40].

The second problem with the use of Cas9 is the off-target cleavage of partially complementary DNA. Errors in DNA recognition may result from inaccurate pairing of guide and target outside of the primary binding region (seed) [40, 41]. There are two fundamental approaches to reducing off-target effects: increasing the specificity of Cas9 [42] and limiting the time of its action on DNA [40, 43–47].

Other representatives of class II CRISPR–Cas systems have also found practical application. In 2015, the effector nuclease Cas12a (also known as Cpf1) from the type V system of bacteria *Acidaminococcus* and *Lachnospiraceae bacterium* was described [48]. Cas12a cleaves both strands of double-stranded DNA with the help of one RuvC nuclease domain, which leads to the formation of a double-strand break with 4–5-nucleotide 5' sticky ends [48, 49]. As in the case of Cas9, the presence of a PAM motif in Cas12a is necessary for recognition of the target, in this case, to
the left of the recognized sequence. After cutting the target DNA and releasing the products, Cas12a can perform non-specific exonuclease cleavage of single-stranded DNA. This phenomenon has been called “collateral activity.” In the cell, such cleavage can disrupt various steps of DNA metabolism. Based on the collateral activity of Cas12a, systems for detecting target DNA sequences in biological samples HOLMES (an one-HOur Low-cost Multipurpose highly Efficient System) [50] and DETECTR (DNA endonuclease-targeted CRISPR trans porter) [51] have been developed.

The type VI effector nuclease of the CRISPR–Cas system, Cas13 (also known as C2c2), has two HEPN (higher eukaryotes and prokaryotes nucleotide binding) RNase domains. This type of nuclease was first isolated in 2016 from the bacterium Leptotrichia shahii [52]. Cas13 recognizes and cleaves single-stranded RNA targets using crRNA, also showing nonspecific collateral activity. Cas13 can be directed to target transcripts in bacterial, plant, and mammalian cells [53]. The efficiency of target RNA cleavage is comparable to the activity of RNA interference using short RNAs (see below). However, Cas13 showed significantly fewer off-target effects. Based on the collateral activity of Cas13, a highly sensitive method for the detection of severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2), the causative agent of COVID-19 [56].

Fig. 2. Cleavage of nucleic acids by Argonaute proteins. Argonaute is loaded with a guide single-stranded nucleic acid molecule (guide binding), followed by a target search based on complementary interactions (target recognition). If the target is complementary to the guide, the conformation of the enzyme changes, catalysis occurs (target cleavage) and the cleaved target is released (target release).

At the end of 2021, two groups of programmable prokaryotic transposon nucleases were described. Transposons of the IS200/IS605 group encode the IcsB proteins, the ancestral forms of Cas9, and TnpB, from which the Cas12 nucleases originated. They can be guided to the target DNA sequence by guide RNA: ơRNA for IcsB and reRNA (right end RNA) for TnpB. As in the case of Cas nucleases, the interaction of these transposon nucleases with target DNA requires the presence of a specific short sequence TAM (target-adjacent motif, or transposon-associated motif) to the right or to the left of the recognized sequence for IcsB and TnpB, respectively. The possibility of introducing targeted changes into the cell genome using these nucleases was shown [57, 58].

ARGONAUTE PROTEINS—ANOTHER GROUP OF PROGRAMMABLE NUCLEASES

Eukaryotic Argonautes—Key RNA Interference Proteins

Argonaute (Ago) proteins, like Cas proteins, recognize and cut the target when loaded with a complementary guide nucleic acid (Fig. 2). Unlike Cas nucleases, no additional sequences in the target (such as a PAM motif) are required for this. Argonaute proteins are found in all domains of life: in many bacteria, archaea, and in almost all eukaryotes, suggesting their ancient origin and functional importance [59–61]. In eukaryotes, Argonaute proteins are involved in the regulation of gene expression by small RNAs, suppression of transposon activity, and protection of cells from viral infections [62–68]. They bind short interfering RNAs (siRNAs), microRNAs (miRNAs), or RNAs associated with the PIWI proteins (piRNAs) and coordinate subsequent gene silencing events usually by interacting with other protein factors [68–71].

The resulting effector complexes recognize complementary target RNAs and can cleave it due to the endonuclease activity of Argonautes. Also, Argonaute-mediated cleavage-independent RNA destabilization and repression of transcription and translation through interactions with other proteins have been described [72, 73].

There are three groups of eukaryotic Argonaute proteins [74–76]. The first includes AGO-like proteins that are involved in cytoplasmic post-transcriptional gene silencing. The second group includes PIWI-like proteins, which are found predominantly in animal germ cells; they suppress the expression of
transposable genetic elements and thus contribute to maintaining the integrity of the genome [69, 77–79]. The third group, WAGO, is represented exclusively by worm proteins and was studied in the Caenorhabditis elegans model. Different types of organisms have different numbers of Argonaute proteins of different groups. For example, four proteins belonging to the AGO group and four to the PIWI group are known in humans [68, 80]. Different RNA interference pathways can operate in the same organism and even in the same cell [81].

RNA interference involving proteins of the AGO group occurs via two alternative pathways, at the level of RNA or chromatin [80, 82–85]. An RNA–protein complex, RISC (RNA induced silencing complex) is responsible for the suppression of gene expression. siRNAs are double-stranded RNAs 20–30 nucleotides long and are formed in the cytoplasm as a result of the processing of exogenous and endogenous double-stranded RNAs by the Dicer enzyme. miRNAs are formed in the nucleus by the Drosha endonuclease from RNA precursors containing a hairpin structure [70, 77, 86–90]. After loading of the Argonaute protein with short double-stranded RNAs, the dissociation of one strand of the RNA duplex occurs, after which such complex with single stranded guide RNA can recognize a fully or partially complementary target mRNA. This results in translation inhibition, endonuclease cleavage of the target by Argonaute, or exonuclease degradation of mRNA by other nucleases [59, 91–93]. RNA interference can also occur at the chromatin level via histone modification or DNA methylation, which also makes it possible to suppress gene expression or, in rare cases, leads to transcription activation [80, 94, 95]. PIWI proteins also translocate into the nucleus upon piRNA loading and suppress transposon expression via transcriptional gene silencing [69, 84, 96, 97]. In general, RNA interference pathways are involved in a wide range of cellular functions, including growth, development, apoptosis, as well as in pathophysiological processes such as carcinogenesis [98].

All eukaryotic Argonautes have four domains: N-terminal, PAZ (Piwi-Argonaute-Zwille), MID (middle) and PIWI (P-element Induced Wimpy Testis), forming a channel in which nucleic acids are located [83, 99–101].

The PIWI domain of Argonaute proteins contains a conserved tetrad of amino acid residues DEDX (where X is an aspartic acid, histidine, or lysine residue), which coordinates two divalent cations required for catalysis [59]. Many Argonaute proteins contain substitutions of negatively charged amino acid residues of the catalytic tetrad and are not capable of target cleavage [59–61].

The MID domain contains a pocket for the 5’ end guide binding. In some cases, the 5’-terminal nucleotide of the guide binds specifically. Thus, human hAgo2, KpAgo from the yeast Kluyveromyces polysporus, and SIWI from the silkworm Bombyx mori bind guides containing an uridine residue at the 5’ end [99, 102, 103]. The corresponding unpaired nucleotide of the target strand can be specifically recognized in a separate pocket of the PIWI domain [104].

PAZ is a small domain (~140 a.a.) that is involved in the binding of the 3’ end of the guide molecule, but is not absolutely necessary for this. Interestingly, PAZ is present not only in Argonautes, but also in the Dicer protein. Probably, the PAZ domain of the ancestor of modern eukaryotic Argonautes underwent a duplication and appeared in Dicer [105].

The N-terminal domain is the least conserved and, possibly, promotes separation of the RNA duplex [99, 106–108].

**Prokaryotic Argonautes: Diversity of Structures**

The existence of Argonaute genes in bacteria and archaea has been known for a long time, and AfAgo from the archaea Archaeoglobus fulgidus [109–112] and AaAgo from the bacterium Aquifex aeolicus [113–115] were even used as models for structural studies of RNA interference. The first description of the diversity of prokaryotic Argonautes was made in 2009 [59]. K. Makarova et al. found 85 homologues of eukaryotic Argonautes encoded in 80 prokaryotic genomes in the non-redundant protein database RefSeq. The rapid growth in the number of sequenced genomes of bacteria and archaea in recent years has made it possible to find many more Argonautes: 487 in 2014 [61], 1010 in 2018 [60], and 1711 in 2020 [116]. In general, about 10–20% of bacterial genomes and about a quarter of archaeal genomes encode at least one Argonaute, and in rare cases, several Argonaute genes at once.

Two large groups can be distinguished on phylogenetic tree of Argonaute proteins (Fig. 3): long Argonautes, containing the same 4 domains as eukaryotic proteins, and short, in which the N-terminal and PAZ domains are absent [59–61]. All short Argonautes have substitutions in the active site in the PIWI domain, which indicates their lack of catalytic activity. Long Argonautes are divided into two groups, called “longA” and “longB” [60]. All representatives of group B are inactive and often have a truncated PAZ domain, while most of the longA Argonautes have a catalytic tetrad and a full-length PAZ. Nevertheless, some representatives of different branches of group A also contain substitutions in the catalytic tetrad, which indicates the multiple appearance of inactive forms in the course of evolution [59–61]. Phylogenetic analysis has shown that eukaryotic Argonautes are a monophyletic group and originate from the longA prokaryotic Argonautes [59–61]. The phylogenetic tree of prokaryotic Argonautes does not correspond to the taxonomic tree of living organisms. From this, it can be concluded that horizontal gene transfer significantly...
contributes to the distribution of Argonaute genes among bacteria and archaea [59–61].

To date, several dozen of prokaryotic Argonautes from different branches of the phylogenetic tree have been characterized. Table 1 describes the properties of most prokaryotic Argonaute proteins studied to date. Attention was mostly focused on in vitro studies of active Argonautes from bacteria and archaea, which have nuclease activity. Unlike eukaryotic Argonautes, which work in the cell with RNA guides and RNA targets, prokaryotic Argonautes preferentially recognize DNA targets. Some of them can also cleave target RNA, although less efficiently; the functional significance of this phenomenon has not yet been elucidated [100, 126, 148]. Most prokaryotic Argonautes bind DNA guides, but some (MpAgo from the bacterium Marinitoga piezophila and its closest homologues) prefer RNA guides [130]. It is known that some eukaryotic Argonautes can also use DNA guides for in vitro target recognition (for example, hAgo2) [107, 156] and even participate in DNA repair (AGO1 from Arabidopsis thaliana) [157], but whether they are capable to interact with DNA in vivo remains unknown.

Unlike eukaryotic Argonautes, prokaryotic Argonautes do not require accessory proteins to be loaded with guide nucleic acids. For TtAgo from Thermus
The MID and PIWI domains are the most conserved and are present in all Argonaute proteins. The MID domain plays a key role in the guide 5’ end binding. In prokaryotic Argonautes and eukaryotic PIWI proteins, the amino acid residues of the MID pocket coordinate a divalent cation (Mg²⁺ or Mn²⁺) involved in the binding of the 5’-terminal phosphate of guide molecule [100, 103, 110, 111, 148, 158]. However, among the active prokaryotic Argonautes, there is a small group of proteins with a hydrophobic pocket in the MID domain. Structural analysis of one of the representatives of this group, MpAgo from Marinibacter piezophila, showed that such Argonautes do not bind the divalent cation in the MID pocket and prefer guides that are not phosphorylated at the 5’ end [60, 130].

Some Argonautes of bacteria and archaea discriminate guides by the nitrogenous base of the 5’-terminal nucleotide: RsAgo from the bacterium Rhodobacter sphaeroides binds guides containing uracil at the 5’ end, TtAgo from the bacterium Thermus thermophilus—cytosine, FpAgo from the archaea Ferroglobus placidus—guanine, MjAgo from archaea Methanocaldococcus jannaschii—purines. At the same time, other studied prokaryotic Argonautes do not show specificity for the 5’-terminal nucleotide (Table 1) [101–103, 122, 124, 125, 130, 141, 142, 146, 159]. The corresponding unpaired nucleotide of the target chain can be specifically recognized in a separate pocket of the PIWI domain (G for TtAgo, A for RsAgo) [145, 158].

Being bound in the MID pocket, the 5’-terminal nucleotide of the guide molecule does not participate in the formation of hydrogen bonds with the complementary nucleotide in target and is called an anchor. There are several regions that can be distinguished in the guide molecule: the primary binding region (“seed”) — 2–8 guide nucleotides, the cleavage site — 10–11 nucleotides, the 3’-supplementary site — 12–16 nucleotides, the 3’-terminal region, and the last nucleotide which usually binds in the pocket of the PAZ domain [151, 160–164]. Most active Argonautes cleave the target between positions complementary to 10 and 11 nucleotides of the guide [100, 101, 132, 133, 148]. Some prokaryotic Argonautes, which normally bind 5’-phosphorylated guides, can also use 5’-OH guides, for example, CbAgo, LrAgo, KmAgo, SeAgo, but in this case the cleavage site is shifted by one nucleotide [122, 123, 126, 142].

Binding of a complementary target by a guide-loaded Argonaute is accompanied by structural rearrangements, which include rotation of the PAZ domain and changes in the position of several loops in the PIWI domain [100, 129, 144, 148]. The guide 3’ end is released from PAZ and a catalytically active ternary Argonaute complex with the guide and target is formed [165, 166].

For eukaryotic Argonautes, guide-target pairing at positions 2–8 of the duplex in the seed region play a crucial role. Non-complementary nucleotides and insertions in the guide opposite the target cleavage site also significantly reduce the efficiency of target cleavage, while mismatches in the 3’ terminal region do not have a significant effect [151, 163, 167]. Such a general pattern cannot be revealed for prokaryotic Argonautes, probably due to their diversity. Interactions of guides with not fully complementary targets may affect the activity of prokaryotic Argonautes in different ways. It has been shown that insertions in the seed region do not affect guide–target duplex binding by TtAgo and RsAgo [158, 168, 169]. In LrAgo and SeAgo, non-complementary interactions in the seed region even stimulate target cleavage [122, 142]. It is interesting to note that the presence of additional or non-complementary nucleotides in the seed region stimulates the release of the guide–target hybrid from hAgo2 and prokaryotic RsAgo, which may be the mechanism for the replacement of Argonaute–associated guide molecules [150, 158]. In contrast to eukaryotic Argonautes, CbAgo, LrAgo, SeAgo, and TtAgo are sensitive to the presence of mismatches in the 3’ supplementary region of the DNA guide [122, 142, 143]. In general, such experiments are necessary to understand the specificity of target binding and cleavage by new prokaryotic Argonautes from different groups.

It is still unknown which structural features of prokaryotic Argonautes are responsible for the discrimination of RNA/DNA guides and targets, as well as for sensitivity to base-pairing in the guide–target duplex. Probably, these features are primarily determined by variations in the structures of the MID and PIWI domains. To answer this question, further studies of the activity and structures of prokaryotic Argonaute proteins are required.

**Argonaute Proteins Protect Prokaryotic Cells from Foreign Nucleic Acids**

A possible role of prokaryotic Argonautes in protecting cells from viruses was suggested in 2009 [59]. At the same time, experimental data confirming this hypothesis have appeared only recently.

First, it has been shown that Argonaute genes are often located in the so-called “defense islands,” regions of the genome with increased mobility, in which various genes are encoded that protect bacteria and archaea from foreign nucleic acids. The best known prokaryotic defense systems are restriction-modification systems and CRISPR–Cas. To date, more than a hundred such systems are known, and this number continues to grow [170–173]. Genes with an
| Host                                | Argonaute, protein ID | Guide, length<sup>a</sup> | 5′ end of the guide<sup>b</sup> | Target          | Reaction temperature<sup>c</sup> | Ion                  | Mismatches in a duplex<sup>d</sup> | Catalytic activity                        | References                  |
|------------------------------------|-----------------------|-----------------------------|---------------------------------|-----------------|-------------------------------|----------------------|-----------------------------------|------------------------------------------|-----------------------------|
| **Prokaryotic Argonaute proteins** |                       |                             |                                 |                 |                               |                      |                                   |                                          |                             |
| *Aquifex aeolicus*                 | AaAgo, WP_010880937.1 | DNA 18–24                   | 5′-P; ?                        | RNA, (DNA ?)    | opt. 55°C; up to 75°C        | Mg<sup>2+</sup>, Mn<sup>2+</sup> | ?                                 | Guide-dependent                        | [113–115]                  |
| *Clostridium butyricum* (strains TK520, CWBI 1009) | ChAgo, WP_058142162.1; CbcAgo, WP_045143632.1 | DNA* (RNA weakly) 16–18 | 5′-P; none | DNA* (RNA weakly) | opt. 37°C; 30–60°C (CbcAgo opt. 37–42°C; 25–55°C) | Mg<sup>2+</sup>, Mn<sup>2+</sup>, Co<sup>2+</sup> | ↓5, 6, 12–15 nt | Guide-dependent; guide-independent | [116, 121–123] |
| *Clostridium perfringens*                   | CpAgo, WP_003477422.1 | DNA 15–30 (≥12)             | 5′-P; none | DNA (RNA)         | opt. 37°C; 4–70°C         | Mg<sup>2+</sup>, Mn<sup>2+</sup> | ↓12–15 nt | Guide-dependent                        | [124]                       |
| *Ferroglobus placidus*                 | FpAgo, WP_012966655.1 | DNA 15–16                   | 5′-P; G1                        | DNA              | 75–99°C                      | Mg<sup>2+</sup>, Mn<sup>2+</sup>, Co<sup>2+</sup>, Ni<sup>2+</sup> | ↑7 nt; ↓3,8,11–15 nt | Guide-dependent                        | [125]                       |
| *Intestinibacter bartlettii*           | IbAgo, WP_007287731.1 | DNA 15–30 (≥14)             | 5′-P; none | DNA              | opt. 37°C; 4–60°C         | Mg<sup>2+</sup>, Mn<sup>2+</sup> | ↓14–16 nt | Guide-dependent                        | [124]                       |
| *Kurthia massiliensis*                | KmAgo, WP_010289662.1 | DNA (RNA) 16–20 (≥12)       | 5′-P; none | DNA (RNA)        | opt. 37°C; 37–60°C; from 25°C | Mn<sup>2+</sup>, Mg<sup>2+</sup>, weakly Co<sup>2+</sup> | ↓4, 5, 10–15 nt (DNA–DNA); ↓4, 8–11 nt (DNA–RNA); ↓3–11 nt (RNA–DNA) | Guide-dependent; guide-independent (37°C) | [126, 127] |
| *Limnothrix rosea*                    | LrAgo, WP_075892274.1 | DNA* 16–18 (≥10)            | 5′-P; none | DNA              | opt. 37°C; 30–54°C        | Mn<sup>2+</sup>, Mg<sup>2+</sup>, weakly Co<sup>2+</sup> | 14–8 nt; ↓10–15 nt | Guide-dependent; guide-independent | [122]                       |
| *Methanocaldococcus fervens*          | MfAgo, WP_015791216.1 | DNA, RNA 16                 | 5′-P DNA and RNA; 5′-OH DNA    | DNA              | opt. 80–90°C; from 54°C    | Mn<sup>2+</sup>, Mg<sup>2+</sup>, Co<sup>2+</sup> | ?                                 | Guide-dependent                        | [128]                       |
| *Marinitoga piezophilia*              | MpAgo, WP_014295921.1 | RNA 16–40                   | 5′-OH; none                     | DNA (RNA)        | 37, 55, 60°C                | Mn<sup>2+</sup>, Mg<sup>2+</sup>, weakly Ni<sup>2+</sup> | ↓5, 7, 8 nt; ↓5–15 nt dinucleotide mismatches | Guide-dependent                        | [129–131]                  |
| Host                          | Argonaute, protein ID | Guide, length | 5'-end of the guide | Target | Reaction temperature | Ion | Mismatches in a duplex | Catalytic activity                  | References |
|-------------------------------|-----------------------|---------------|---------------------|--------|----------------------|-----|------------------------|-------------------------------------|------------|
| **Methanothermobacterium jannaschii** | MjAgo, WP_010870838.1 | DNA 15–41     | 5'-P; purines       | DNA    | opt. 75–95°C; from 37°C | Mg$^{2+}$ | ?                      | Guide-dependent; guide-independent | [101, 132, 133] |
| **Natronobacterium gregoryi**  | NgAgo, WP_005580376.1 | DNA 22–24     | 5'-P; ?             | DNA    | 37°C                 | Mg$^{2+}$ | ?                      | Guide-dependent                    | [134–138] |
| **Pyrococcus furiosus**       | PfAgo, WP_011011654.1 | DNA 15–31     | 5'-P; none          | DNA    | opt. 87–99.9°C; from 37°C | Mn$^{2+}$, Co$^{2+}$ | ?                     | Guide-dependent; guide-independent | [139–141] |
| **Synechoccocus elongatus**   | SeAgo, WP_011244830.1 | DNA* 14–24    | 5'-P; none          | DNA*   | opt. 37°C             | Mn$^{2+}$, weakly Mg$^{2+}$ | ↑5, 6, 10 nt; ↓7, 12–15, 18 nt | Guide-dependent; guide-independent | [142] |
| **Thermotoga profundula**    | TpAgo, WP_041081268.1 | RNA 21        | 5'-OH; ?            | DNA    | 60°C                 | Mg$^{2+}$, Mn$^{2+}$ | ?                      | Guide-dependent                    | [130] |
| **Thermus thermophilus**      | TtAgo, WP_011174533.1 | DNA* 13–25    | 5'-P; Cl1 G1'       | DNA*   | opt. 73–75°C; ≥20°C ssDNA; ≥65°C plasmids | Mn$^{2+}$, Mg$^{2+}$ | ↓12–15 nt        | Guide-independent                  | [100, 106, 143–149] |
| **Eukaryotic Argonaute proteins** |                        |               |                     |        |                      |     |                       |                                     |            |
| **Homo sapiens**              | hAgo2, NP_036286.2    | RNA* (DNA) 20–21 | 5'-P; U1 A1'        | RNA*   | 37°C                 | Mg$^{2+}$ | ↓2–8 nt (target binding); ↓8–11 triple mismatch | Guide-dependent                    | [99, 139, 150–152] |
| **Kluyveromyces polysporus**  | KpAgo, XP_001644461.1 | RNA* (DNA) 12–17 (11–25) | 5'-P; U1 | RNA* | 25–30°C | Mg$^{2+}$ | ↓7–14 nt (↓9–13 nt) | Guide-dependent                    | [102, 153] |
| **Bombyx mori**               | SIWI, NP_001098066.2  | RNA* 28–30    | 5'-P; U1            | RNA*   | 26, 37°C             | Mg$^{2+}$, Mn$^{2+}$ | ↓2–7 nt (target binding) | Guide-dependent                    | [103, 154, 155] |

*aThe type of guide nucleic acid and its length (number of nucleotides).

bCharacteristics of the 5'-end of the guide: P—phosphate group, OH—hydroxyl group; the presence of specificity to the nitrogenous base is indicated: none—no specificity, A/U/G/C—types of nitrogenous bases, 1—the first nucleotide of the guide chain; 1’—the first nucleotide of the target chain. Here and further: ?—unknown.

cThe temperature at which Argonaute protein could cleave target; opt.—the temperature optimum, the temperature range at which the Argonaute cleaves targets is also given. Discrete temperature values are given for publications where measurements were carried out only under these conditions.

dThe effect of mismatches (unpaired nucleotides of the duplex guide–target) on the efficiency of target cleavage; the positions of the nucleotides (nt) are numbered relative to the 5'-end of the guide. ↑—increase in cleavage efficiency, ↓—decrease in cleavage efficiency.

eBoth in vitro and in vivo.
unknown function, often encoded in defense islands, are also likely to be involved in defense systems [59, 174–176]. Interestingly, some groups of prokaryotic Argonautes are associated with certain defense systems. For example, MpAgo, which uses RNA guides, and its closest homologues are encoded inside CRISPR-Cas systems [130]. A more extensive analysis showed that active prokaryotic Argonautes in most cases are encoded in genomes that also contain CRISPR-Cas systems, while genomes without Argonautes or with inactive Argonautes contain CRISPR-Cas systems much less often [116]. A more detailed analysis of the association of various groups of Argonautes with other defense systems of prokaryotes is the subject of further research.

Secondly, some prokaryotic Argonautes are associated with additional domains that perform protective functions. Thus, many short Argonautes contain the SIR2 (silent information regulator 2) or TIR (Toll/interleukin-1 receptor) domains or their coding sequences are located in the neighboring gene [59, 60]. These domains are also part of the Thoeris defense system found in bacteria: in response to infection of a cell with a virus, they cleave NAD+, which leads to cell death and stops the spread of the virus in the population [175, 177, 178]. It has recently been shown that the SIR2/Ago and TIR/Ago systems protect bacterial cells from phages and plasmids: short Argonautes with MID-PIWI domains recognize foreign DNA, while SIR2 or TIR cleaves NAD+ [178, 179]. The TIR/Ago system is also involved in antiplasmid defense. Some Argonautes are associated in operons with nucleases from different families, which are presumably involved in the production of guides or act as protein effectors of defense systems [59, 60].

Finally, direct experimental confirmation of the protective role of long Argonautes in bacteria has been obtained in recent years. Thus, the CbAgo protein protects cells from phages [116], the CbAgo [116], PfAgo (from the archaea Pyrococcus furiosus) [141], and RsAgo [159] proteins reduce the efficiency of transformation and promote plasmid degradation in bacterial cells.

In addition to protecting cells from foreign DNA, Argonautes can perform other functions. To date, one such example is known: it has been shown that, in the absence of gyrase, TtAgo is involved in cell division, resolving chromosome catenanes after replication in Thermus thermophilus cells [149]. The diversity of the structures and activities of prokaryotic Argonautes may indicate that their functional role in bacterial and archaean cells is not limited to currently known examples.

**PROSPECTS FOR THE PRACTICAL APPLICATION OF PROKARYOTIC ARGONAUTE PROTEINS**

Argonaute proteins are considered promising candidates for directed nucleic acid manipulation and genome editing. The ability of Argonaute proteins to act as site-specific programmable nucleases opens up wide opportunities for their use in vitro. The currently known applications of bacterial Argonaute proteins are summarized in Fig. 4.

Unlike restriction endonucleases, Argonaute proteins can be used to make a cut in potentially any DNA sequence and obtain fragments with “sticky” ends of the required length and nucleotide composition. Since Argonautes have a single active site and can cleave only one strand of DNA, double-stranded DNA cleavage requires the loading of Argonautes with a pair of guides that are complementary to opposite strands at the cutting site. For example, PfAgo from a thermophilic archaea was successfully used to assemble genetic constructs and to analyze them [185]. In a large-scale study using the guide panel for the TtAgo, patterns of preference for guide sequences were determined. Based on the results obtained in the course of this work, it is possible to optimize the using of TtAgo as a programmable endonuclease [143, 186].

An important area of application of Argonaute proteins is the detection of target sequences and nucleotide modifications in biological samples. A system for detecting the desired sequence in the sample was developed on the basis of PfAgo called PAND (PfAgo-mediated Nucleic acid Detection). The technique uses reporter DNA, upon cleavage of which by Argonaute, the fluorescent label and quencher are uncoupled and a fluorescent signal appears (Fig. 4). The guide for the reporter cleavage is produced from the target DNA sequence and is cut from the target DNA of the sample by PfAgo using a combination of three synthetic guides. When using reporters with different fluorescent beacons, it is possible to detect several sequences in one tube [180]. This method has been successfully adapted to identify SARS-CoV-2 RNA and its mutant variant (D614G in the spike protein) in patient samples [187]. Combining two methods, PAND and ligase chain reaction made it possible to simplify the detection of nucleic acids. The combined protocol was called PLCR (PfAgo coupled with modified Ligase Chain Reaction for nucleic acid detection). A thermostable ligase joins two halves of the guide using the target DNA as a template. Then, in the course of several cycles, guide is amplified. The guide is loaded into the Argonaute, and the reporter DNA is cut, which leads to appearance of fluorescent signal [181].

Other proposed methods are based on the decreased activity of Argonautes in the presence of mismatches or modifications in the target, compared to the situation when the guide and the target are completely complementary (Fig. 4). A quick analysis of the
presence of insertions or deletions in one or two alleles of the target gene that occurs during genome editing is possible if guides are selected to the editing nuclease cleavage site. In the case of the wild-type allele, the DNA will be cut by the Argonaute. The reaction products can be detected in a gel [182]. The NAVIGATER method (Nucleic Acids of clinical interest Via DNA-Guided Argonaute from *Thermus thermophilus*) has been developed to detect rare single nucleotide substitutions that occur, for example, in cancer cells. To detect DNA or RNA with a single nucleotide substitution, it is necessary to choose a guide that is complementary to the predominant wild-type sequences in the sample, and a rare variant has a substitution at the 10th or 11th position of the target, counting from the 5' end of the guide. The presence of the substitution dramatically reduces the nuclease activity of TtAgo. In this case, the wild-type DNA will be cleaved by the Argonaute, and the relative amount of the rare variant will increase. Such enrichment makes it possible to detect the rare variant by standard methods (various PCR techniques or sequencing) [183].

To detect substitutions and modifications in RNA, it is necessary to use Argonaute with the appropriate specificity. MpAgo uses RNA as guides and can bind complementary target RNA. For target recognition, nucleotides 6 and 7 are most important. By determining the dissociation constant of the guide RNA with the target, it is possible to identify mismatches, as well as modified nucleotides such as inosine (Fig. 4) [131]. The difference in the cleavage efficiency of single- and double-stranded RNA can be used to study the structure of highly structured RNA. The ability to guide the Argonaute to specific RNA sequences makes it possi-
Argonautes also found application in super-resolution microscopy to visualize structures beyond the diffraction limit of microscopy. The DNA-PAINT (DNA Point Accumulation In Nanoscale Topology) method is based on the detection of association and dissociation of a fluorescent DNA probe with a fixed DNA target. If labelled DNA is loaded into CbAgo, the rate of the process increases due to the fact that Argonaute organizes guide DNA for optimal target binding [188]. The ability of Argonaute to accelerate the binding of a fluorescently labeled DNA guide to target RNA was also used for specific visualization of miRNA. The authors, S. Shin et al. [184], called this technology Ago–FISH (Argonaute-based Fluorescence In Situ Hybridization) (Fig. 4).

The ability of Argonautes to introduce site-specific cuts in nucleic acids allows us to consider them as potential tools for editing the genome and transcriptome. As such, Argonautes may have some advantages over Cas nucleases. First, Argonautes do not require the presence of a PAM sequence in the target DNA. Second, the guides for most prokaryotic Argonautes are short DNA molecules while Cas nucleases require long RNA guides. DNA synthesis is much cheaper, which can facilitate development of the Argonaute-based genome editing system. Third, the size of the Argonaute molecule is smaller than that of Cas9, so it is easier to deliver them inside cells [189, 190].

In 2016, a work was published on editing the human cell genome using Argonaute from the halophilic archaea Natronobacterium gregoryi (NgAgo) [189]. However, the authors soon retracted the article, since other groups of researchers did not confirm the reliability of the results presented in it when trying to introduce changes using NgAgo in the genome of human cells, mouse embryos and Danio rerio embryos, as well as the hepatitis B virus [135, 137, 138, 191–194]. An attempt to edit the genome of human cells using Argonaute TtAgo also failed [135]. Thus, the question of the possibility of using Argonautes as a tool for editing the genome remains open. It should be noted that the choice of the Argonaute protein can significantly affect the results of such experiments. In published works, Argonautes from therophilic microorganisms were used, which have an optimum efficiency at high temperatures. In addition, NgAgo is a protein from the halophilic archaea, and it is known that the expression and folding of such proteins occurs inefficiently under normal salt conditions [195]. In some studies, which did not give a positive result, a nuclear localization signal (NLS) was added to the C-terminus of the Argonaute [135, 137, 193]. This could affect the activity of the protein, since the C-terminus of Argonaute is important for catalysis [144]. Guides to sequences located on opposite DNA strands at a great distance from each other were also used [194]. Several studies using NgAgo have noticed a decrease in the mRNA level of the target gene without making changes to the corresponding genome region, possibly due to Argonaute interactions with target RNAs [137, 138]. Thus, for further experiments on genome editing, it is worth to choose Argonautes highly specific for DNA targets, whose activity is optimal under physiological conditions.

Argonautes with a non-functional catalytic center, inactive Argonautes or mutant forms of active Argonautes, also have great potential for practical applications. Such Argonautes recognize target DNA or RNA sequences using guides and can potentially be used as a DNA recognition domain in the construction of chimeric proteins with various functions. Previously, similar approaches were successfully used in the case of Cas nucleases. Possible applications of catalytically inactive bacterial Argonaute proteins are summarized in Fig. 5. For example, the combination of Argonautes with enzymes that modify nitrogenous bases may be used to make changes in the DNA or RNA sequence without creating a double-strand break. Cytidine deaminase converts cytidine to uridine, which leads to the replacement of the C:G pair with T:A during DNA replication [196]. A reverse substitution of the T:A pair with C:G can be carried out by a chimeric protein containing adenosine deaminase as a functional domain, which converts adenine to inosine, which is recognized by polymerases as guanine [197].

Argonautes may be used as a DNA recognition domain to recruit factors that change epigenetic marks and thus activate or inactivate gene expression (Fig. 5). For this, approaches previously proposed for catalytically inactive variants of Cas nucleases can be used. Activation or suppression of gene expression may be achieved by combining Argonaute with transcriptional activators or repressors, respectively. For example, directing cytosine-DNA-methyltransferase DNMT3a to the promoter of the target gene can reduce its expression. The use of the KRAB domain (Krüppel-associated box), which forms a complex with two histone DNA methyltransferases, leads to the repression of target genes. Fusion with p300 acetyltransferase leads to acetylation of lysine at position 27 of histone H3 and, as a result, transcriptional activation of target genes [40, 198]. Changes in the arrangement of DNA regions in the nucleus can lead to changes in gene expression, for example, when promoters and enhancers become closer to each other [199]. For this, inactive Argonautes with heterodimerizing protein domains can be used. Argonautes may be also used as a DNA recognition domain for visualizing various structures and processes in the nucleus when fused with fluorescent proteins, as was shown for TALEN nucleases [200]. Also Argonautes could be used as an RNA recognition domain for visualizing the location of target RNA in the cell (Fig. 5). Inactive Argonautes interacting with RNA targets could be used to attract additional functional domains to RNA to regulate gene expression at
the post-transcriptional level. For example, the effector domains of the YTHDF1 and YTHDF2 proteins (YT521-B Homology Domain Family) in eukaryotic cells recognize N6-methyladenosine in RNA. YTHDF1 activates translation of the target RNA, while YTHDF2, on the contrary, leads to its degradation. Replacement of the N6-methyladenosine-recognizing domain with an RNA-recognizing domain of guide loaded Argonaut makes it possible to direct these functions to any RNA [201]. It is also possible to perform directed RNA editing using the ADAR1 or human ADAR2 deaminase domain (Adenosine Deaminase Acting on RNA). This protein converts adenosine to inosine, mainly in cases where adenosine is opposite cytidine in the RNA duplex [202]. Similar to published experiments using dCas9 (a catalytically inactive variant of Cas-9), RNA-recognizing Argonauts could also be used to regulate alternative splicing by changing the efficiency of incorporation of certain exons into mature mRNA (Fig. 5) [203].

The tools for genome editing has evolved from the search and creation of proteins capable of recognizing specific DNA sequences (meganucleases, TALEN, zinc finger nucleases) to the discovery of universal programmable nucleases, the specificity of which is determined by guide nucleic acids. These targeted nucleases include enzymes based on Cas proteins, which are currently actively used for gene editing, as well as Argonauts proteins. Although no success was yet achieved in the use of Argonauts for genome editing, their diversity, both structural and functional, allows us to hope that they will soon provide complementary tools for genome manipulations and biotechnology along with Cas nucleases.

At present, studies of the prokaryotic Argonate family proteins already led to their practical applications in new highly sensitive methods for detecting nucleic acids, including SARS-CoV-2 RNA. In the near future, we could expect the appearance of works using Argonauts as effective tools for manipulations with nucleic acids in vitro, and in vivo.

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The authors declare that they have no conflicts of interest. This article does not contain any studies involving animals or human participants performed by any of the authors.
AUTHOR CONTRIBUTIONS

E.V. Kropocheva, L.A. Lisitskaya, A.A. Agapov, and D.M. Esyunina contributed equally to this work.

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