AsnI: a novel class II restriction endonuclease from Arthrobacter sp., strain N-CM, recognizing 5'-AT/TAAT-3'

Bernhard U. Rexer, Michael Jarsch, Christine Sagmeister, Barbara Glück, Gerhard Berger and Christoph Kessler*

Boehringer Mannheim GmbH, Biochemical Research Center, Department of Molecular Biology, Nonnenwald 2, D-8133-Penzberg, FRG

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A new class II restriction endonuclease, AsnI, with a novel sequence specificity was isolated from the Gram-positive eubacterium Arthrobacter species, strain N-CM. AsnI recognizes the unambiguously defined palindromic hexanucleotide

consisting of A- and T-residues. The novel enzyme in the presence of Mg²⁺ cleaves specifically both strands as indicated by the arrows. The staggered cuts generate 5'-protruding ends with single-stranded 5'-TA-3' dinucleotide extensions. The novel enzyme may be a useful tool for cloning experiments by complementation of the few enzymes such as *PstI* and *PvuI* cutting only once in the Amp^r-gene of plasmids pBR322 and pBR328.

Restriction endonuclease; Hexanucleotide recognition sequence; T/A-nucleotide; 5'-Protruding terminus; Klenow enzyme; T₄ polynucleotide kinase; T₄ DNA polymerase; T₄DNA ligase

1. INTRODUCTION

More than 650 class II restriction endonucleases have been isolated from eu- and archaebacteria [1,2]. These enzymes represent more than 100 different sequence specificities. Most important are enzymes recognizing unequivocally defined tetra-and hexanucleotide palindromes. Out of the 16 possible tetranucleotides, 11 sequences are currently covered by corresponding enzymes.

From the 64 possible hexanucleotides, 49 enzyme specificities are known so far [3]. Most of these enzymes recognize sequences rich in G- and C-residues. With SspI and DraI as well as its isoschizomer AhaIII only tree enzymes are known

Correspondence address: C. Kessler, Boehringer Mannheim GmbH, Abtlg BB-BRM, Nonnenwald 2, 8122 Penzberg, FRG

recognizing hexanucleotide palindromes consisting only of A- and T-residues [4-6].

We have discovered a further enzyme, AsnI, recognizing only A- and T-residues within the hexanucleotide palindrome 5'-AT/TAAT-3'. We report here the isolation and characterization of the novel enzyme enlarging the set of class II restriction endonucleases cutting at AT-rich hexanucleotides.

We designated the new class II enzyme in accordance with the proposal of Smith and Nathans [7] as AsnI to distinguish it from the series of enzymes isolated from different strains of Achromobacter species [1].

2. MATERIALS AND METHODS

2.1. Bacterial strain and culture conditions
Arthrobacter sp., strain N-CM, belongs to the irregular, non-

sporing Gram-positive rods [8]. Cells were cultured aerobically at 25°C in a mixture of 10 g/ml malt extract and 10 g/l yeast extract at a pH value between 6.8 and 7.0.

2.2. Enzymes

Restriction endonucleases Asp7001, Aval, BamHI, BglI, EcoRI, EcoRV, HindII, NdeI, PstI, PvuI, PvuII, SalI and ScaI, calf intestinal alkaline phosphatase (CIAP), T₄ DNA polymerase (T₄ DNAP), T₄ DNA ligase (T₄ DNALig), T₄ polynucleotide kinase (T₄ PNK), terminal transferase (TdT) and Klenow enzyme (PolIK) were from Boehringer Mannheim. The enzymes were employed according to the manufacturer's specifications.

2.3. DNAs, nucleotides and reagents

Phage λ c1857Sam7 and M13mpl8RF DNA, plasmids pUC18, pBR322 and pBR328, DNA molecular mass markers III and IV, dATP, dGTP, dCTP and dTTP were from Boehringer Mannheim. Ad-2 DNA was from BRL. [γ -32P]ATP (\sim 3000 Ci/mmol) and [α -32P]ddATP (\sim 3000 Ci/mmol) were from Amersham, heparin-Sepharose CL-6B, Sephadex G25 from P-L Pharmacia, and cellulose PII was from Whatman.

2.4. Isolation and purification of AsnI

Asnl was purified according to standard procedures [3] including fractination on Sephadex G25, cellulose Pll and heparin-Sepharose CL-6B.

2.5. Enzyme assay

For determination of AsnI activity, varying amounts of the final enzyme preparation were incubated with $1 \mu g \lambda c 1857 Sam7$ DNA in 25 μl incubation mixture (10 mM Tris-HCl, pH 8.0/37°C; 100 mM NaCl; 10 mM β -mercaptoethanol; 10 mM MgCl₂) for 1 h at 37°C. The reactions were terminated by addition of 5 μl cold stop solution. 25 μl of each reaction mixture was resolved by electrophoresis as above.

One unit of Asn1 is defined as the amount of enzyme which digests 1 μg of $\lambda c1857Sam7$ DNA within 1 h at 37°C under the stated assay conditions.

2.6. DNA labelling

Restriction fragments were 5'-endlabelled by treatment with 0.05 units of CIAP per pmol 5'-termini [9] and rephosphorylation with 1 unit T₄ PNK per pmol 5'-termini in the presence of $50\,\mu\text{Ci}$ [γ -\$^2P]ATP (~3000 Ci/mmol) [10]. 3'-endlabelling was achieved with 25 units TdT per pmol 3'-termini and 50 μ Ci [α -\$^2P]ddATP (~3000 Ci/mmol) as described by [11].

2.7. Determination of cleavage positions

Determination of cleavage positions within both strands of AsnI cleavage sites using the chemical method was according to [12] and [13]. Nucleotide sequence analysis was performed following the chemical method described in [10]. The A/G-reaction was modified according to [14]. Treatment with T₄ DNAP was performed as described in [13].

2.8. Computer analysis

Computer data were generated on a Microvax II (Digital Equipment Corporation) under VAX/VMS using the UWGCG software package [23]. In addition, computer programs were used which are analogous to those described in [24]. Sequence

data for Ad-2 DNA [15] and for $\lambda c 1857 Sam7$ DNA [16] were obtained from the EMBL databank [17], Entries AD2 and LAMBDA. The additional nucleotide sequences used for computer analysis were determined by [18,22] for pUC18 DNA and M13mp18RF DNA, by [19,20,22] for pBR322 DNA, and by [21,22] for pBR328 DNA.

3. RESULTS

3.1. Optimal conditions for enzyme activity

The activity of AsnI is optimal to pH 8.0 (37°C) and salt concentrations of 100 mM NaCl. The enzyme is strictly dependent on Mg²⁺ with an optimum at 10 mM MgCl₂, but does not require S-adenosyl-methionine or ATP for activity.

Highly purified AsnI is free of contaminating site-specific or unspecific endonucleases as well as exonucleases, since either $\lambda c1857Sam7$ DNA or pBR322 DNA digested with at least 20 units AsnI for 16 h (320-fold excess of enzyme) gave sharp bands without any smearing after electrophoresis in agarose gels (not shown). Functional purity is shown by ligation with 0.1 units T₄ DNALig per μg AsnI-fragments which yields in >95% ligation products >95% of the ligation products can be recut with AsnI.

3.2. Determination of recognition sequence

The recognition sequence of AsnI was determined by mapping its recognition sites on pBR322 DNA at position 3539, on pBR328 DNA at position 2450, and on pUC18 DNA at positions 516, 635 and 1870 by double digestions with AsnI and various restriction enzymes cutting around these sites.

With pBR322 after linearization of the plasmid with AsnI, double digestions were performed with a series of restriction endonucleases including BamHI, BglI, EcoRV, HindIII, NdeI, PstI, PvuI, PvuII and SalI (fig.1). For all tested enzymes, the resulting fragment sizes are in agreement with AsnI recognition at the hexonucleotide 5'-ATTAAT-3' sequence at position 3539 (table 1). The position of the single AsnI recognition site on pBR328 DNA at position 2450 was determined in an analogous way with PstI (pos. 2520), PvuI (pos. 2647) and Asp700I (pos. 2874) double digestions.

In addition, the three AsnI recognition sites on pUC18 DNA at positions 576, 635 and 1870 were mapped with the help of BglI (pos. 245, 1813),

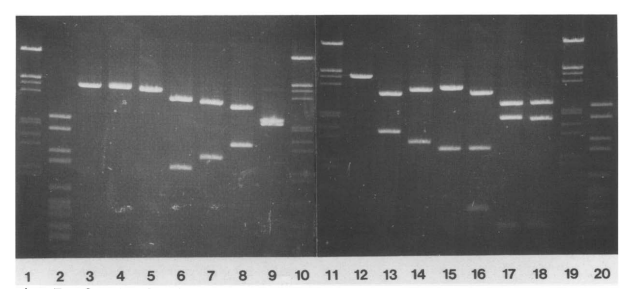


Fig. 1. Double digestions of pBR322 DNA with AsnI and additional restriction enzymes. pBR322 was cut with AsnI (lanes 3, 12), and the resulting linearized DNA was additionally cleaved with PstI (lane 4), PvuI (lane 5), EcoRV (lane 6), BamHI (lane 7), SalI (lane 8), AvaI (lane 9), PvuII (lane 13), NdeI (lane 14), HindII (lane 16), and BglI (lane 18), respectively. pBR322-HindII fragments (3256, 1107 bp) are in lane 15; pBR322-BglI fragments (2319, 1810, 234 bp) are in lane 17. λc1857Sam7-EcoRI/HindIII fragments (lanes 1, 10, 11, 19), and pBR328-BglI/HinfI fragments (lanes 2, 20) were used as DNA molecular mass markers.

PstI (pos. 411), EcoRI (pos. 450) and PvuI (pos. 276, 2066) (not shown). The cleavage sites on pUC18 DNA were localized independently by

Table 1

Mapping of the single AsnI recognition sequence on pBR322

DNA at position 3539 by double digestions with additional enzymes

Second enzyme		Experimentally observed fragment sizes of double digestions (bp)	Computer-derived fragment sizes of double digestions (bp)
PstI	3609	4300, (<100) ^c	4290, 73
PvuI	3736	4200, 200	4165, 198
EcoRV	187	3400, 1000	3351, 1012
BamHl	375	3200, 1200	3165, 1198
SalI	651	2900, 1500	2889, 1474
AvaI	1425	2200, 2100	2248, 2115
PvuII	2066	2900, 1500	2891, 1472
NdeI	2296	3100, 1200	3120, 1243
HindII	651, 3907 ^a	2900, 1100, 350	2887, 1107, 369
Bg/I	929, 1163,	2300, 1700, 250,	2319, 1758, 234
	3482 ^b	(<100) ^d	52

^aResulting fragment sizes: 3256, 1107 bp

determination of the exact cut positions within the respective recognition sequences at positions 577, 636 and 1871.

The AsnI recognition sequence 5'-ATTAAT-3' was confirmed by comparison of the experimentally observed AsnI fragment pattern obtained by digestion of λcI857Sam7 DNA at recognition sites 3418, 19 481, 23 083, 23 692, 24 986, 26 762, 27 202, 30 619, 33 221, 33 735, 33 800, 34 686, 36 274, 43 409, 46 407, 46 852 and 48 377, Ad-2 DNA at positions 31 900, 33 079, 35 425 and 35 936 and M13mp18RF DNA at 4131, 4135, 4238, 4628, 6046, 6105 and 6966 with the corresponding computer data derived by analysis of the known nucleotide sequences with the program MAP of the UWGCG program package [23].

3.3. Determination of the cleavage positions

The exact cleavage position of AsnI was determined in parallel by two approaches, the chemical [10,12] and enzymatic sequencing method [13]. First, for chemical sequencing ³²P-endlabelled DNA fragments containing AsnI sites were prepared from plasmid pUC18. Four aliquots of each of these fragments were subjected to the various chemical sequencing reactions. Another

^bResulting fragment sizes: 2319, 1810, 234 bp

^cDecrease of 4363 bp fragment size

^dDecrease of 1810 bp fragment size.

aliquot was treated with AsnI. After electrophoretic separation on 5% (w/v) polyacrylamide gels, containing 8 M urea, and autoradiography, the size of the AsnI fragments was analysed by comparison with the sequencing ladders as shown in fig.2.

In detail, the cut positions within the AsnI recognition sequence were determined as follows: plasmid pUC18 was linearized by PvuI. This fragment was labelled at the 3'-ends with TdT and $[\alpha^{-32}P]ddATP$. After a second cleavage with EcoRI a 1.6 kb fragment was isolated after electrophoretic separation on an agarose gel. In a second experiment pUC18 DNA was linearized by PstI. The resulting fragment was 3'-endlabelled with TdT and $[\alpha^{-32}P]ddATP$, or 5'-endlabelled with T₄ PNK and $[\gamma^{-32}P]ATP$. After recutting with

PvuI, 1.6 kb fragments were isolated after electrophoretic separation on an agrose gel. In the next experiment pUC18 DNA was linearized by EcoRI. The resulting fragment was either 3'-endlabelled with TdT and $[\alpha^{-32}P]ddATP$, or 5'-endlabelled with T₄ PNK and $[\gamma^{-32}P]ATP$. After recutting with PvuI 1.6 kb fragments were isolated after electrophoretic separation on an agarose gel.

Each of these isolated endlabelled DNA fragments were divided into five aliquots. Four of these aliquots are subjected to base-specific cleavage reactions according to [12]. One aliquot was treated with AsnI. Reactions products were analyzed by electrophoresis on 5% (w/v) polyacrylamide gels, containing 8 M urea, and subsequent autoradiography. Interpretation of the results followed the rules described in [13].

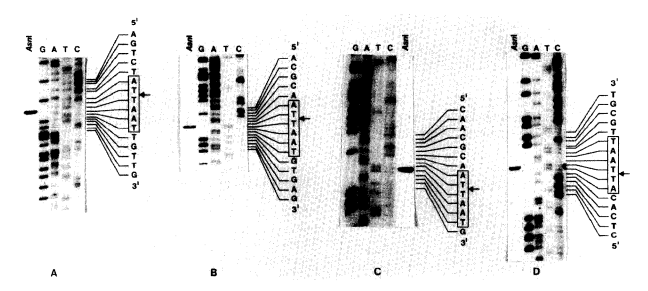


Fig. 2. Autoradiograph of the sequencing ladders around AsnI recognition sites on plasmid pUC18 at positions 1870 (A) and 576 (B,C,D). To analyze the AsnI site at position 1870 of pUC18 DNA, the plasmid was cut with PvuI, subsequently 3'-endlabelled with TdT and $[\alpha^{-32}P]$ ddATP, and recut with EcoRI. An endlabelled PvuI-EcoRI fragment (A) was isolated by low melting temperature agarose electrophoresis. For the analysis of the Asn1 site at position 576 pUC18 DNA was either cut with Pst1, 3'-endlabelled with TdT and $[\alpha^{-32}P]$ ddATP, or cut with EcoRI, either 5'-endlabelled with T₄ PNK and $[\gamma^{-32}P]$ ATP, or 3'-endlabelled with TdT and [α-32P]ddATP. All samples were recut with PvuI. For the sequencing ladder a suitable 3'-endlabelled PstI-PvuI fragment (B), a 3'-endlabelled EcoRI-PvuI fragment (C), and a 5'-endlabelled EcoRI-PvuI fragment (D) was prepared as above. The isolated fragments were divided into aliquots which were subjected to chemical sequencing reactions according to [10] with the modification of the A+G reaction described in [14]. An additional aliquot of each fragment was treated with AsnI. The resulting samples were analyzed by electrophoresis on 5% (w/v) polyacrylamide sequencing gels, containing 8 M urea, followed by autoradiography. Lanes G, A, T, C represent the four chemical sequencing reactions of a particular DNA fragment, lane AsnI represents the respective AsnItreated sample. The interpretation followed the rules described in [13]. The cut sites of Asn1 at positions 1871 (A) and 577 (B,C,D) were deduced from the size of the corresponding bands of the sequencing ladder. During the base-specific chemical sequencing reactions a particular base is eliminated leaving a phosphate group at both flanking 5'- and 3'-DNA termini. In the case of the 5'-endlabelled sample (D) this results in a slightly faster electrophoretic mobility as compared with the corresponding enzymatically created fragment which has lost only its 3'-terminal phosphate group leaving 5'-phosphorylated fragment termini [26].

Cleavage positions of AsnI on pUC18 DNA determined by chemical sequencing are as follows:

The specificity of AsnI is concluded as:

These results were confirmed by the enzymatic sequencing approach described in [13]. A M13mp18-derivative containing an insert with an AsnI site was used as template for enzymatic sequencing reactions starting with the universal sequencing primer. In parallel, an universal sequencing primer, ³²P-endlabelled with T₄ PNK and $[\alpha^{-32}P]ATP$, was annealed to the same template, and a partial double-stranded DNA was created by treatment with PolIK and all four dNTPs. This double-stranded DNA was used as substrate for AsnI to produce a 5'-endlabelled DNA fragment comparable to the sequencing ladder. Samples were analyzed by electrophoresis and subsequent autoradiography as described above (fig.3). The cleavage position of AsnI determined by enzymatic sequencing on M13 DNA is as follows:

*5'---CCGGTAT TAATTCATG-3'

Again, the specificity of AsnI is concluded as:

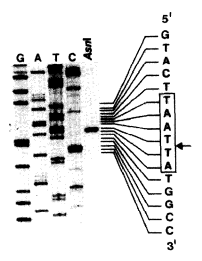


Fig. 3. Autoradiograph of the sequencing ladder around the AsnI recognition site of M13mp18 DNA. Preparation of single-stranded M13mp18 DNA and sequencing reactions were done according to [27]. A partial double-stranded, 5'-endlabelled DNA as substrate for AsnI was prepared and PolIK extension of a 5'-endlabelled universal primer annealed to single-stranded M13mp18 DNA. Lanes G, A, T, C represent the four base-specific sequencing reactions, lane AsnI represents the respective AsnI-treated sample. The interpretation followed the rules described in [13]. The fragment resulting from AsnI cleavage has the identical size compared with the band of the first T in the sequencing ladder of the AsnI recognition sequence ATTAAT. This band of the T sequence ladder represents a fragment ending with 5'...AT-3', and therefore implicates the

cleavage mode: AT TAAT.

4. DISCUSSION

With AsnI from Arthrobacter species, strain N-CM, a new class II restriction endonuclease has been isolated in high purity recognizing the novel palindromic hexanucleotide recognition sequence 5'-AT/TAAT-3'.

The new enzyme complements class II restriction endonucleases recognizing AT-rich hexanucleotide sequences. Out of the eight possible hexanucleotide palindromes existing completely of A-and T-residues, corresponding enzymes for only two sequences 5'-AATATT-3' (SspI [4]) and 5'-TTTAAA-3' (DraI [5], AhaIII [6]) were known to date. In contrast, for most of recognition sequences rich in G- and C-residues enzymes have been discovered. For the eight possible hexa-

nucleotide palindromes composed completely of G- and C-residues many more class II enzymes have been discovered so far. Only the 5'-CGCGCG-3' sequence has not yet been realized by a class II restriction endonuclease.

AsnI may be a useful tool for the construction of recombinant DNA. The single AsnI cleavage site in the commonly used cloning vectors pBR322 and pBR328 is located in the structural gene for ampicillin resistance. Cloning into the unique AsnI site should be facilitated because in contrast to single-cutting enzymes PvuI, ScaI or Asp700I, creating blunt ends, AsnI generates fragments with 5'-protruding AT-dinucleotides. Compared with the 3'-protruding fragment ends obtained by linearization of pBR322 of pBR328 DNA with PstI, AsnI fragment termini can both be efficiently 5'-labelled with T4 polynucleotide kinase and 3'-endlabelled with either Klenow enzyme or T4 DNA polymerase.

Finally, as with SspI, DraI and its isoschizomer AhaIII, AsnI may be applied for analysis of ATrich DNA sequence stretches, or as a rare cutting class II restriction endonuclease for genomes of high G:C-content.

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