Full genome sequence of a novel circo-like virus detected in an adult European eel (Anguilla anguilla) showing signs of cauliflower disease

Andor Doszpoly1*, Zoltán L. Tarján1, Róbert Glávits2, Tamás Müller3, Mária Benkő4

1 Institute for Veterinary Medical Research, Centre for Agricultural Research, Hungarian Academy of Sciences, Budapest, Hungary
2 Veterinary Diagnostic Directorate, National Food Chain Safety Office, Budapest, Hungary
3 Department of Fish Culture, Szent István University, Gödöllő, Hungary

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Author’s address: Andor Doszpoly, Institute for Veterinary Medical Research, Centre for Agricultural Research, Hungarian Academy of Sciences, H-1581 Budapest, P.O. Box 18, Hungary; e-mail: adoszpoly@vmri.hu
Abstract

An adult European eel (Anguilla anguilla), showing typical signs of the so-called cauliflower disease, was subjected to pathological and molecular virological examinations. Samples, taken from the internal organs and the polypoid proliferative tissue from the mouth, were examined by PCR for the detection of several viruses. Positive results were obtained with a nested PCR targeting the rep gene of circoviruses. Analysis of the partial rep sequence indicated the presence of a putative novel circovirus, but attempts to isolate it remained unsuccessful. The missing part of the genome was acquired by an inverse nested PCR with two specific primer pairs, designed from the newly determined rep sequence, then genome walking was applied. The circular full genome was found to consist of 1378 nt. Two oppositely oriented ORFs were present. One of them could be identified as a circoviral rep gene unambiguously. However, the predicted product of the other ORF, though it is a clear positional counterpart of the cap genes, showed no obvious homology to any known circoviral capsid proteins. A stem-loop-like element in the intergenic region between the 5’ ends of the ORFs was also found. Phylogenetic calculations indicated that the novel virus belongs to the Circovirus genus of the Circoviridae family. The relative amount of the viral DNA in the organ samples was estimated by quantitative real-time PCR. The results suggested that the examined fish was caught in an active viraemic status, albeit the role of this circovirus in the etiology of the cauliflower diseases could not be ascertained.
Introduction

The cauliflower disease (stomatopapilloma or orocutaneous papillomatosis) of the European eel (*Anguilla anguilla*) was first described at the beginning of the 20th century (Wolf 1988). Sporadic occurrence of the disease, with no specificity for age and size of the affected fish, has been reported mostly from the tributary regions of northern European rivers (Delves-Broughton et al. 1980; Wolf 1988). The benign epidermal neoplasms usually grow on the head region, but lesions may occur on other parts of the body (Peters 1975). The papillomas consist of fibro-epithelium and are sometimes pigmented. The cause of the tissue proliferation is unknown. Different viruses, resembling birnav-, orthomyxo- and rhabdoviruses have been detected in or isolated from eels with cauliflower disease, but experimental reproduction of the clinical signs has not been successful (Nagabayashi & Wolf 1979; Schwanz-Pfitzer et al. 1984; Ahne & Thomsen, 1985; Ahne et al. 1987). Because of the seasonality of the tumor development, the effect of chemical pollutants and water temperature changes were assumed to contribute to the manifestation of the disease (Peters 1975).

The European eel, presently a critically endangered species, were being introduced to Lake Balaton (the largest lake in Central Europe, located in the western part of Hungary) regularly from 1961 to 1991, except in 1985 and between 1989 and 1990. As a result, more than 83 million elvers (glass eel stage, which start to feed) were stocked in Lake Balaton. By the end of the 1980s, the lake had become overpopulated with eel, and there were two massive periods of eel deaths. These deaths were probably due to a nematode worm (*Anguillicoloides crassus*) infection in 1991 and 1995 (Molnár et al. 1991). Thereafter the Hungarian Government prohibited the introduction of additional eels. By now, the eel stock in Lake Balaton is composed of very old individuals, of which even the youngest are more than 23 years old.
In this work, the detection and full genome analysis of a novel circo-like virus are described. The virus was found in different organs of an eel from Lake Balaton which exhibited typical signs of the so-called cauliflower disease.

Circoviruses (CVs) are small, naked DNA viruses. The icosahedral virions, with a diameter of 12—26 nm, contain a circular single-stranded DNA genome. The genome size of the CVs studied to date was found to range between 1.7-2.3 kb. The viral capsid is composed of one structural protein (capsid protein) (Biagini et al. 2011). Targeted screenings with a sensitive consensus nested PCR (Halami et al. 2008) resulted in the recognition of a large and still sharply increasing number of novel CVs and circo-like viruses in an incredible diversity of specimens including natural and waste waters, fecal samples from different mammals from bats to man, as well as in several invertebrates and on the surface of a variety of raw meat products (Blinkova et al. 2009; Li et al. 2010; Li et al. 2011). Up until a few years ago, only birds and swine had been known as frequent hosts for CVs. In the past several years, the vertebrate host spectrum has widened to include fish and amphibians (Lőrincz et al. 2011 & 2012; Tarján et al. 2014). However, in the majority of these cases, the eventual pathological role of the detected CVs is unknown.

Material and methods

Origin of the sample. During a routine limnological survey, an approximately 50-cm-long adult eel with cauliflower-like growths on both sides of its mouth was caught in Lake Balaton, Hungary on 21st September 2009. After euthanasia, a necropsy was performed. Small pieces from the labial tissue proliferation, the gills, heart, liver, spleen and kidney were collected in duplicate. One part of the samples was fixed in Bouin’s fixative, embedded in paraffin, sectioned (at 4—5 µm), stained with hematoxylin and eosin, and viewed by light microscopy.
according to standard procedures. The other set of samples were frozen, and used for virus isolation and molecular studies.

**Virus isolation.** Virus isolation was attempted on the EK-1 cell line (Chen et al. 1982). The pooled organ homogenates were diluted to a 10% (w/v) suspension in an L-15 medium (Gibco) complemented with antibiotics (Penicillin 300 U/ml, Streptomycin 300 µg/ml). The suspension was centrifuged at 2000 × g for 10 min, and the supernatant was transferred into a new tube immediately. Three parallel inoculations (500 µl suspension per flask) were made in 25-cm² flasks of EK-1 monolayers at 80% confluency. The flasks were incubated at three temperatures (15°C, 19°C, and 22°C) and checked for the appearance of cytopathic effect (CPE) daily.

**PCR.** For PCR, the nucleic acid extraction was made from approximately 25 mg tissue as described in detail by Dán et al. (2003). The tissues were homogenized in a 1× TE buffer, and 100 µl from each organ suspension was digested with proteinase K (20 mg/ml) in the presence of sarcosyl (10%). After incubation with guanidine-hydrochloride (8 M) for one hour at room temperature, the DNA was precipitated with ethanol. The presence of adenoviral DNA in the organ samples was tested by a sensitive nested PCR with consensus primers targeting the DNA dependent DNA polymerase gene (Wellehan et al. 2004). For the detection of herpes- irido- or poxviral DNA, a broad spectrum PCR method was used (Hanson et al. 2006). Demonstration of the circoviral DNA was attempted by a widely used and very efficient consensus nested PCR described by Halami et al. (2008). For the amplification of the missing part of the putative circular genome of the newly detected CV, two specific primer pairs were designed and used in a nested inverse PCR. The primer sequences were as follows:

outer forward: 5'-GCG CTT GAG GAT TCT CAT TC-3'; outer reverse: 5'-CAG ATC GTT CCT CTT CCC TT-3'; inner forward: 5'-GAC TTT GGA TGG AAG AAG CC-3'; inner reverse: 5'-CCT TGT TAT GCT GGT CGT TG-3'.
The PCR program consisted of an initial denaturation step of 98°C for 5 min, followed by 45 cycles of 98°C for 30 sec/56°C for 30 sec/72°C for 60 sec and a final elongation cycle of 72°C for 3 min. The reaction mixture consisted of 34 µl distilled water, 10 µl of 5×HF buffer (Phusion, Thermo Scientific), 0.5 µl Phusion enzyme (Phusion, Thermo Scientific), 1 µl (50 µM) of each (forward and reverse) primer, 1.5 µl of dNTP solution of 10 mM concentration, and 2 µl of the target DNA in a final volume of 50 µl. The reactions were performed in a T1 Thermocycler (Biometra). The results of the PCRs were analyzed by electrophoresis in agarose gels.

Quantitative, real-time PCRs for determining the relative amount of the viral DNA in five organs (the labial proliferative tissue, the gills, liver, spleen and kidney) were carried out in an Applied Biosystems® StepOnePlus™ Real-Time PCR System instrument (Life Technologies). Two specific primers, suitable for the amplification of a 100-bp product were designed. The reaction mixture contained 25 µl 2× PrimeSTAR Max Premix (Takara Bio Inc.), 17.5 µl distilled water, 2.5 µl EvaGreen™ Dye (Biotium) 1 µl of each primer (forward: 5'-AGGCAACGACCAGCATAACA-3'; reverse: 5'-AGTCGTCGATGCAGGCCAAG-3') and 3 µl of the target DNA in a final volume of 50 µl. The program consisted of an initial denaturing at 98°C for 5 min, followed by 40 cycles of 98°C for 10 sec, 55°C for 5 sec, and 72°C for 10 sec. The beta-actin gene (with forward 5'-ACCGGTATCGTCATGGACTC-3'; and reverse 5'-CGTCAGGGTCTTTCATCAGGT-3’ primers) was used as an internal standard. The results were analyzed by the StepOne Software v2.1 (Applied Biosystems).

Sequencing and sequence analyses. All PCR products were excised from the gels, purified with the QIAquick Gel Extraction Kit (Qiagen), and sequenced directly with the inner primers. The sequencing reactions were performed with the use of the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems). The electrophoresis was carried out by a commercial service provider on an ABI PRISM 3100 Genetic Analyzer. The amplification
product from the inverse PCR, encompassing more than 1000 bp, was cloned with the use of
the CloneJET Kit (Fermentas), and sequenced with the primers supplied with the kit. The
newly obtained nucleotide sequences were compared with their homologs in the GenBank by
using different BLAST algorithms at the NCBI portal. The genome was assembled manually
and confirmed with the use of the Staden Package as described elsewhere recently (Doszpoly
& Shchelkunov 2010). The size and orientation of ORFs and putative genes were examined
after a 6-frame translation of the genomic DNA with the use of the JavaScript DNA
Translator 1.1 program (Perry 2003).

**Phylogeny inference.** Phylogenetic calculations were performed online at the Mobyle portal
(http://mobyle.pasteur.fr/cgi-bin/portal.py) of the Pasteur Institute (Paris) using the distance
matrix analysis (Protdist) with the Jones–Taylor–Thornton matrix, then the Fitch calculations
were performed with global rearrangements. The tree topology was tested by bootstrap
analysis (Seqboot/1000 samplings, Protdist, Fitch, Consense).

**Results**

**Gross and Microscopic Pathology.** On both sides of the mouth, soft polypoid, raspberry-
coloured masses (approximately 1 cm in diameter) were seen (Fig 1a). The fish was slightly
emaciated. The only other gross lesions were restricted to the heart where a small flat grayish
oval-shaped patch (of about 5 mm in diameter) was revealed on the epicardial surface which
extended into the myocardium (Fig. 2a). By light microscopy, no lesions were seen in the gills
or kidney. The labial polypoid masses were benign papillomas composed of proliferating
connective tissue and covered by a multilayered epithelium. Proliferation of the malpighian
cells upon a narrow base of connective tissue without an invasion down into the underlying
dermis was also seen (Fig. 1b). The grayish spot on the heart was a benign myoma composed
of collections of cross-striated muscle fibres and surrounded by connective tissue (Fig. 2b).
Additionally, vacuolization in the hepatocytes, and multifocal hemosiderin deposition in the red pulp of the spleen were revealed.

**Virus isolation.** Rounding and detachment of the cells appeared on 12, 14 and 17 days post-inoculation in the tissue cultures incubated at 22°C, 19°C and 15°C, respectively. In the negative controls, no CPE was observed. A few days later, the CPE was complete. However, in the second and third passages, no CPE was observed. PCR testing of samples from the successive passages for the detection of the CV or certain dsDNA viruses (adeno-, herpes- and iridovirus) also remained negative.

**PCR and sequencing.** All PCRs, performed for the detection of different large DNA viruses, gave negative results. However, the circovirus PCR resulted in specific products from every examined organ (including the gills, liver, spleen and kidney) as well as from the labial proliferative tissue. Interestingly, the gills, the spleen, and the labial tissue all initially displayed a heavy longitudinal smear and discrete bands of amplicons were obtained after only 10× dilution of the target DNA solutions. The size of the identical amplicons was found to be 303 bp after editing out the primer sequences. By homology search, the conservative region of the circoviral replication-associated protein (Rep) was identified. The inverse nested PCR yielded an 1192 bp DNA fragment. It was first sequenced directly, then it was cloned and sequenced again for a better quality outcome. This fragment contained the whole *cap* gene, the rest of the *rep* gene with the two intergenic regions.

**Genome sequence analysis.** The complete genome of the putative eel CV was found to encompass 1378 nt with an average G+C content of 48.7%. The full genome sequence was deposited to the GenBank and assigned to accession number KC469701. The genome organization was found to be somewhat divergent from one typical of CVs, inasmuch as the putative *cap* gene proved to be significantly shorter (Fig. 3). Nonetheless, two oppositely
oriented major ORFs which flanked a putative stem-loop element were found. The stem-loop element, situated in the 5’ intergenic region, possessed 12-bp stem. Its loop region consisted of 12 nucleotides with the conserved circoviral nonamer sequence (TAGTATTAC) therein as shown in Fig. 4. The deduced product of the rep gene was predicted to consist of 286 amino acid (aa) residues containing the conserved RNA helicase domain. Its closest homologue in the GenBank, with 51% aa identity, was the corresponding region of the CV recently described in the wels catfish (Lőrincz et al. 2012). The other ORF, supposedly corresponding to the gene of the capsid protein (Cap), was predicted to code for 114 aa only. This protein did not show convincing homology to any proteins in the GenBank. Nonetheless, similar to the Cap proteins of other known CVs, a 32 aa-long arginine-rich stretch (RRRYRRKSNRPRNCQRRYRRPRRERNR) was found close to the N terminus. The length of the intergenic regions between these ORFs was 107 and 65 nt, at their 5’ and 3’ termini, respectively. Two additional, oppositely oriented ORFs overlapping the rep gene were also found. One of these, in the same orientation as the rep gene, encompasses 309 nt, the translated aa sequence shows homology to a small hypothetical (17 kDa) protein of the beak and feather disease virus (Niagro et al. 1998). The other ORF, in a reverse position compared to rep, consists of 483 nt. Its deduced product shows homology to hypothetical genes described from pigeon and duck CVs (Mankertz et al. 2000; Chen et al. 2006). It has to be mentioned however, that this 483-nt ORF has no ATG. A possible alternative start codon could be the ATA triplet (in nt position 19 to 21). If so, then the predicted protein product would consist of 155 aa.

**Quantitative PCR.** With qPCR, the virus was successfully detected in all five of the examined organs. The statistical mean of five parallel qPCRs for the viral genome resulted in the following cycle threshold (Ct) values: liver 23.6, spleen 23.9, gills 26.0, kidney 27.5, stomatopapilloma 28.1. It means that the relative amount of the viral DNA in the examined
organs is the following: 1 unit in the tissues of the stomatopapilloma, 1.25 unit in the kidney, 20 units in the gills, 130 units in the spleen, and 157 units in the liver.

Phylogenetic analysis. The phylogeny reconstruction made by the distance matrix analysis on complete deduced Rep sequences from 27 CVs is presented in Fig. 5. The alignments consisted of 229 aa. The newly detected putative European eel CV appeared in the clade of the Circovirus genus within the Circoviridae family as a sister group of the wels catfish CV (Lőrincz et al. 2012). However, the monophyly of fish CVs was not confirmed.

Discussion

In the present work, the full genome sequence of a putative novel CV was determined. The virus originated from a European eel showing typical signs of the so-called cauliflower disease. We propose to name this virus the European eel circovirus (EeCV). The isolation of the virus on an eel kidney cell line failed. The CPE, observed after the first inoculation, did not appear again in the consecutive passages. We speculate that the cell degeneration was probably due to a direct cytotoxic effect by some material or unknown, uncharacterized virus that was not detectable by our PCRs.

Apart from the heart myoma and the stomatopapillomas, which were probably large enough to interfere with feeding, no other significant pathological alterations were revealed. It seems that the examined eel was caught in a viraemic status, since every examined organ contained circoviral DNA. The initial PCR with undiluted samples of the stomatopapilloma, gills, and spleen resulted in a heavy smear. This signaled the possibility of too high concentration of the total DNA. Indeed, the 10-fold dilution of the same samples produced discrete bands of the expected size.
Thanks to the significant improvement in the efficiency of detection methods (Halami et al. 2008), an incredible increase in the number of the known small circular ssDNA viruses has occurred in the second half of the past decade. The known host range of CVs (including mainly birds and swine) has also grown rapidly (Delwart & Li, 2012). Besides a couple of additional mammals, a number of invertebrate hosts have also been found to harbor seemingly specific CVs. The first CVs from lower vertebrates have been published most recently. These viruses, detected in barbel (*Barbus barbus*) and wels catfish (*Silurus glanis*), have been characterized by full genome analyses (Lőrincz et al. 2011 & 2012). Additional putative piscine CVs have been discovered by PCR in other fishes, as well as in two species of amphibians. But, these CVs have not been confirmed yet by complete genomic sequences (Fehér et al. 2013; Tarján et al. 2014). Although the barbel and wels catfish CVs have been found during investigations for the cause of increased mortality, the role of these viruses in disease could not be confirmed.

In this case, a direct connection between the presence of the putative circoviral DNA and the cauliflower disease could be revealed. The porcine and most avian CVs are known to have an immunosuppressive effect (Todd 2004), which may exacerbate the pathogenicity of certain concurrent infectious agents. Nonetheless, it is also possible that CVs can cause generalized infection in individuals whose immune system is temporarily or permanently impaired. Ng et al. (2009) have drawn similar conclusions when they examined a novel small ssDNA virus in sea turtles with fibropapillomas.

The taxonomy of EeCV is somewhat ambiguous. According to the phylogeny reconstruction based on the Rep sequence, the piscine (barbel and wels catfish) CVs seem to form a common clade with an interesting small circular ssDNA virus. This common clade is formed with NG13, which was recently discovered by screening human fecal samples collected in Nigeria (Li et al. 2010). The NG13 virus had originally been described as an
outlier to the proposed new genus *Cyclovirus* (Li et al. 2011) however, its exact taxonomic place is still unclear. The nonamer sequence of the stem-loop of the NG13 is identical to that of the CVs, but its genome organization resembles that of the cycloviruses (CyVs) (Delwart & Li, 2012). Our analysis shows that NG13 and fish CVs might share a close evolutionary origin. It seems that the EeCV does not form a monophyletic group with the other two fish CVs. A possible reason for such a result could be the incomplete taxon representation. Perhaps the full genome analyses of additional novel CVs from fishes and other lower vertebrates will help discern the phylogenetic relationships more precisely in the future.

The family *Circoviridae* is currently facing a radical revision. It has been proposed that the *Gyrovirus* genus be moved to the *Anelloviridae* family. Furthermore, the establishment of a new genus, *Cyclovirus* within the *Circoviridae* has also been proposed (Biagini et al. 2013).

There are however, numerous circovirus-like viruses with unknown ancestry at the moment.

Based on its genome organization, the putative EeCV most closely resembled the members of the *Circovirus* genus. It encodes a Rep protein on the virus sense strand and a putative Cap protein in the opposite direction. The presence of the two additional smaller ORFs exhibit a clear homology to the hypothetical genes of yet unknown function, described in pigeon (Mankertz et al. 2000) and duck CVs (Chen et al. 2006), and in the beak and feather disease virus (Niagro et al. 1998). These homologies further confirm the place of EeCV in the same genus. Furthermore, the stem-loop structure, which is involved in the initiation of the viral genome replication (Steinfeldt et al. 2001), had a typical circoviral nonamer sequence in EeCV, identical to that of the fish CVs, porcine CV1, pigeon, gull and finch CVs (Li et al. 2010). The lengths of the intergenic regions between the *rep* and *cap* genes also show differences between CVs and CyVs (Li et al. 2010). These regions in EeCV are also characteristic for CVs. Its closest evolutionary relative is the wels catfish CV.
Interestingly, the exceptionally short cap gene of EeCV did not show obvious homology to its counterparts in any known CVs or CyVs. Yet, a characteristic feature of the circoviral Cap proteins, namely the arginine-rich stretch close to the N-terminus, was identified.

It is intriguing to speculate about a recombination event between a progenitor and an unknown circular virus resulting in EeCV. Among ssDNA viruses (even originating from distantly related virus families), recombination events occur frequently (Gibbs & Weiller, 1999; Martin et al. 2011). Moreover, a recombination between a circo-like virus and ssRNA virus has also been hypothesized (Diemer & Stedman, 2012). At the moment however, the origin of the cap gene of EeCV remains enigmatic since the BLAST searches did not find any reliable homology to any known sequences among the environmental samples.

According to the proposed new rules of the Circoviridae Study Group of the ICTV, the comparative analysis of the complete genomes would replace the earlier practice of comparing the cap sequences in the establishment of new species (Biagini et al. 2013). Considering the demarcation criteria, EeCV unequivocally represents a novel species.

At the discussion of the qPCR results, we emphasize that these results are based on the examination of one specimen and at one time-point. Thus, these data should be considered as preliminary results. The qPCR confirmed the results of the conventional PCR, inasmuch as all examined organs contained the viral DNA suggesting that the examined eel was caught in a viraemic status. The relative amount of the viral DNA may indicate that the viral replication takes place in the liver and spleen.

To examine the pathogenicity of EeCV in experimental infections, the isolation of the virus would be essential. Nonetheless, screening eels, especially those showing clinical signs of the cauliflower disease, for the presence of cicoviral DNA should also be continued.

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FIGURE LEGENDS

Figure 1a. Anguilla anguilla. Clinical appearance and microstructure of the cauliflower disease. (A) Bilateral labial polipoid masses in the eel from the Lake Balaton. (B) Microphotograph of the labial mass composed of proliferating connective tissue covered by multilayer epithelium. Proliferation of the malpighian cells upon a narrow base of connective tissue without invasion down into the underlying dermis.
**Figure 2a.** Macroscopic picture of the abdominal cavity. The white arrow points to the lesion on the epicardium. (B) Microphotograph of the benign myoma composed of collections of cross-striated muscle fibres and surrounded by connective tissue.

**Figure 4.** Genome organization of the EeCV. Besides the two major ORFs, there are two additional hypothetical ORFs. One of them (BKFDV ORF5) shows similarity to the ORF 5 of Beak and feather disease virus (BKFDV). The other (DuCV and PiCV) is homologous to hypothetical ORFs of duck (DuCV) and pigeon (PiCV) CVs.
Figure 4. The structure and sequence of the putative stem-loop
Figure. 5. Phylogenetic tree reconstruction based on the distance matrix (JTT model) of the deduced amino acid sequences (229 aa) of the rep genes of ssDNA viruses. Bootstrap values are shown on the branches. The cycloviruses, the circoviruses and the fish circoviruses within the family Circoviridae are designated by different colored lines on the tree. Abbreviations: GuCV=gull circovirus; PiCV=pigeon circovirus; StCV=starling circovirus; FiCV=finch circovirus; RaCV=raven circovirus; CaCV=canary circovirus; BKFD-1=beak and feather disease virus 1; BKFD-2=beak and feather disease virus 2; MDuCV=muscovy duck circovirus; MuCV=mallard circovirus; SwCV=swan circovirus; GoCV=goose circovirus; CanCV=canine circovirus; PCV-1=porcine circovirus 1; PCV-2= porcine circovirus 2; NG13=human stool associated circovirus; BaCV=barbel circovirus; CfCV=catfish circovirus; EeCV=European eel circovirus; Fwc= Florida woods cockroach-associated cyclovirus; Df-5=dragonfly cyclovirus 5; GF-4c=bat cyclovirus; Chimp11=chimpanzee Cyclovirus 11; Df-4=dragonfly Cyclovirus 4 ; Df-1=dragonfly cyclovirus 1; Bt-5=bat feces cyclovirus ; TN25=human feces cyclovirus
Pepper golden mosaic virus

GuCV
PiCV
StCV
FiCV
RaCV
CaCV
BKFD-1
BKFD-2
MDuCV
MuCV
SwCV
GoCV
CanCV
PCV-1
PCV-2

NG13
BaCV
CfCV
EeCV

Fwc
df-5
GF-4c
Chimp11
df-4
df-1
Bt-5
TN25

464

0.2

465