Multiple Oct2 isoforms are generated by alternative splicing

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ABSTRACT

The interaction of the Oct2 transcription factor with the cognate octamer motif ATGCAAAT is a critical determinant of the lymphoid-specific expression of immunoglobulin genes. Ectopic expression of cloned Oct2 cDNA was shown to be sufficient to reconstitute at least some aspects of this regulation in non-lymphoid cells. We describe the isolation and characterization of multiple cDNAs encoding mouse Oct2 from a mature B-cell line and we show that a variety of isoforms of this transcription factor is generated from a single gene by an alternative splicing mechanism. All the isoforms retain the previously characterized POU-domain and are therefore able to bind to the octamer motif. Different amounts of the various isoforms are present within the same B-cell regardless of the developmental stage of B-cell differentiation and at least some of the isoforms are conserved between mouse and humans. In cotransfection experiments we show that all the isoforms are able to activate an octamer containing promoter element in fibroblasts revealing an unexpected functional redundancy. Finally, we show that one of the isoforms encodes the previously described lymphoid-specific Oct2B protein which has been suggested to be involved in the function of the octamer motif in the context of the immunoglobulin heavy-chain (IgH) enhancer.

INTRODUCTION

The lymphoid-specific expression of immunoglobulin genes is regulated by at least two distinct cis-acting regulatory elements, the upstream promoter and the intronic enhancer element (1, 2, 3, 4). The octamer motif ATGCAAAT or its reverse complement ATTTGCAT is conserved both in the intronic heavy-chain enhancer as well as in all heavy and light-chain gene promoters (5, 6). It contributes to the lymphoid-specific activity of the enhancer (7) and plays a pivotal role for immunoglobulin promoter activity (8, 9, 10).

However, the octamer motif is not exclusively associated with regulatory elements involved in lymphoid-specific expression. It has also been found to play an important role in the regulation of ubiquitously expressed genes. Cell cycle regulation of histone H2B gene expression has been shown to be governed by the same octamer motif (11). Transcription of genes encoding several of the small nuclear RNAs (U1snRNAs) by both RNA polymerase II and III is also dependent on a functional upstream octamer motif (12, 13). Furthermore, the octamer element has also been implicated in adenovirus replication (14).

Several proteins that interact with the conserved octamer motif have been identified and characterized to different extents. Oct1 (other names are NF-A1, OTF1, OBP 100, NFIII) is a 90-100kD protein which seems to be ubiquitously expressed (15, 16, 17, 18). Two lymphoid-specific octamer binding proteins have been described: Oct2A (other designations OTF2, NF-A2), a protein of 55–60kD (19, 20, 21) and Oct2B (OTF2B) a 75kD protein that has been shown by a proteolytic clipping electrophoretic mobility shift assay to be related to Oct2A (22). Oct4 (NF-A3, Oct3) has been detected as a protein specifically expressed in undifferentiated embryonal stem cells and embryonal carcinoma cell lines (7, 23, 24). In addition, various other octamer binding proteins have been identified in extracts from different cell types and tissues (23, 25, 26).

Recently, cDNA clones were obtained for Oct1, Oct2A and Oct4 (17, 24, 27, 28, 29, 30, 31). Analysis of these clones revealed that they are members of a novel class of homeo-box containing transcription factors characterized by the presence of a POU-domain. The POU-domain was originally described as a region of homology conserved in the mammalian transcription factors Pit-1, Oct-1 and Oct-2 as well as the Caenorhabditis elegans unc-86 gene. It consists of a POU-specific (75–80 amino acids long) and a POU-homeo (60 amino acids long) subdomain separated by a short linker (17, 32). cDNAs encoding additional POU-domain proteins have been isolated using a PCR cloning protocol (33). Results of site directed mutagenesis and deletion experiments suggested that this POU-domain in its entirety is responsible for the sequence-specific DNA-binding as well as for protein-protein interactions (34, 35, 36, 37).

Comparison of the independently isolated human Oct2A cDNA

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sequences revealed that the cDNAs isolated from the BJA-B cell line encode a protein that contains an insert of 16 amino acids that is absent from the protein encoded by the cDNA clone isolated from Namalwa cells (28, 29, 38). It was unclear whether the two forms originated by transcription of highly related genes or whether the difference was due to the presence of two alleles or an alternative splicing mechanism. Here we report the isolation and characterization of multiple functional isoforms of Oct2 cDNA from murine B-cells. We show that they are generated by an alternative splicing mechanism. Among the isolated isoforms are both homologues of the two described human cDNAs and a clone encoding Oct2B.

MATERIALS AND METHODS

Isolation of cDNA clones

The mouse pre B-cell library (kind gift of Dr. P. Nielson, MPI Freiburg) in agt11 was independently screened with two end-labelled 61 base long oligonucleotides. These oligonucleotides were homologous to the published human Oct2A sequence from position 889 to 950 and from 997 to 1058, respectively (28). For PCR amplification the following two primer oligonucleotides were used:

Oct 5': CTGGGATCCGGCAGCATGGTCGACTCC-AGCATGGGCGTCCTAGC
Oct 3': AGGAATTCGAGGGGAGGCATGGCCGGCTCTC-ACTCAGCTTG

The 5' primer contains a BamHI cloning site upstream and a Sall site downstream of the AUG start codon, the 3' primer contains an EcoRI cloning site. First strand cDNA was prepared from 2 μg of total cytoplasmic WEHI231 RNA using oligo(dT) primers and MuLV Reverse Transcriptase (Gibco-BRL). 10% of this cDNA was directly PCR amplified in a final volume of 30 μl using 1 unit of Taq-polymerase (Perkin Elmer) for 40 cycles of 40 seconds at 94°C, 3 min 72°C. The amplified DNA was precipitated, digested with EcoRI and BamHI and cloned into the BluescriptSK+ vector. 90 clones containing Oct2 sequences were selected for further analyses. DNA was digested with MstI which yields about 15–20 fragments that could be resolved on 40 cm polyacrylamide gels. This analysis allowed classification of 70 of the clones into the classes described (Oct2.1–Oct2.6). The remaining 20 clones were single isolates and have not been characterized further.

Isolation of genomic clones

Complete Oct2 cDNA inserts were used for screening of a mouse 129 strain cosmids library (kind gift of A. Poustka) and a single cosmid was repeatedly isolated. As this cosmid was lacking the first 3 coding exons, we constructed a λEMBL3 genomic library from C57Bl/6 mouse spleen DNA. 2 x 10⁶ phages from the unamplified library were screened using a probe containing the first 180bp of the Oct2 cDNA and two phages containing non-overlapping inserts were isolated. One of them was contiguous with the previously isolated cosmid clone and contained exons 2 and 3. The other started about 2 kb upstream of the first phase insert (determined by genomic Southern blots with several enzymes) and contained exon 1.

Subcloning and sequencing

Fragments from the genomic clones were subcloned into the pBluescriptSK+ vector. For sequence analysis multiple additional subclones were generated for deoxy sequencing with M13 universal/reverse primers. In addition several oligonucleotides were generated from the Oct2 cDNA sequence and used for sequencing. Sequencing was done on double stranded DNA using dITP as well as standard nucleotides and T7 DNA Polymerase (Pharmacia) according to manufacturers specifications.

In vitro transcription and translation, electrophoretic mobility shift assays (EMSA)

cDNA containing plasmids (1 μg) were linearized with EcoRI and RNA was synthesized using T3 RNA polymerase according to manufacturers instructions (Stratagene). In vitro translation with 1 μg of RNA was performed in rabbit reticulocyte lysates (Promega) following manufacturers specifications. 10μCi 35S-labelled methionine was included in the reactions. 1/25 of the in vitro translation was analyzed on a 10% SDS polyacrylamide gel. As probes for the EMSA, wild-type or mutant octamer oligonucleotides cloned into the BamHI site of pUC18 and subsequently excised with EcoRI/HindIII were used (19). Binding reactions contained 50 mM NaCl, 10 mM Tris/HCl pH 7.5, 1 mM DTT, 1 mM EDTA, 5% glycerol, 3 μg poly(dI-dC), 5000 cpm of the respective probe and 1/25 of the in vitro translation reaction. Native gels were run in 0.5 x TBE buffer, dried and exposed for 12 hrs with two layers of paper interleaved to suppress signals originating from the labelled proteins.

Cell and tissue culture and transfections

Lymphoid cells were maintained in IScove's modified MEM supplemented with 5% fetal calf serum, 10 μM β-mercaptoethanol and antibiotics. NIH/3T3 cells were grown in DMEM supplemented with 5% calf serum, 5% newborn calf serum and antibiotics. For mitogen stimulations spleen single cells (about 5 x 10⁶ cells/ml) were stimulated with either 30 μg/ml lipopolysaccharide (Sigma) or with 3 μg/ml concanavalinA (Sigma) for 48 hrs. Transfections were done by the calcium-phosphate co-precipitation method. 2 x 10⁶ cells were transfected with 5 μg of the reporter plasmids plus 2 μg of the Oct2 cDNA expression constructs. Sheared salmon sperm DNA was added to adjust the amount of transfected DNA to 20 μg. 4 hrs after the addition of the DNA precipitate a DMSO shock was performed. Cytoplasmic RNA was prepared 44 hrs later as described (39).

RNA analysis

S1 analysis with the mouse histone H4 probe was performed as described (10). For primer extension analysis 30mg of cytoplasmic RNA was co-precipitated with 0.01pmol (typically about 25,000 cpm) of 5' end labelled primer ('5'GGGGTTGTGAGTCCAGATCGATCTG3'). Hybridization was performed in 10μl of 250mM KCl, 10 μM Tris/HCl pH 7.5, 1 mM EDTA for 1 hr at 60°C, then 40μl of 1 x Reverse Transcriptase buffer containing 0.5 mM of each dNTP, 0.1 mg BSA and 200 U MuLV Reverse Transcriptase (Gibco-BRL) was added. Reactions were terminated after 1 hr at 37°C, precipitated and analyzed on 10% denaturing polyacrylamide gels. RNase protection analysis was essentially performed as described (40). RNA from primary cells and organs was isolated.
by the guanidinium-isothiocyanate/CsCl centrifugation method (40). Cytoplasmic RNA from cell lines was prepared as described (39). Typically 10 to 30 μg RNA was hybridized to 500,000 cpm of RNA probe synthesized by transcription with either T3 or T7 RNA polymerase. Protected fragments were analyzed on 6% denaturing polyacrylamide gels.

**Western analysis**

150 μg of nuclear proteins prepared as described (41) were separated on a 10% SDS polyacrylamide gel (40) and then electrophoretically transferred for 2 hrs at 80 V to a nitrocellulose membrane using a BioRad transblot cell. The membrane was stained with PonceauS to visualize the molecular weight standards and was blocked using 2% dry milk after destaining. The blots were reacted with mouse antisera raised against bacterially produced Oct2 isoforms (42) and subsequently with an alkaline phosphatase coupled goat anti-mouse antibody (Jackson laboratories). The blots were developed as described (42).

**RESULTS**

Multiple Oct2 related RNA species are present in B-cells
Initially we had isolated a partial Oct2 cDNA clone by standard techniques from a murine pre B-cell library using oligonucleotides homologous to the POU-specific region and the POU-homeodomain of the published human Oct2A cDNA sequence as probes (38). This clone did not extend to the amino terminus of the coding region (data not shown). We subsequently used oligonucleotide primers from within this partial cDNA clone and primers that were either homologous to the amino terminal

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**Figure 1:** Nucleotide and protein sequences of the various Oct2 isoforms. A.) The composite nucleotide sequence of the mouse cDNAs is shown. The sequence starts 8 or 23 bp upstream of the AUG start codon, depending on which one of the two in frame AUG codons is used (AUG codons underlined). The sequence of the first 40 bp is actually taken from the genomic clones because the oligonucleotide used for PCR amplification contained several point mutations in order to introduce convenient cloning sites. The regions encoding the POU-specific domain and the POU-homeo domain are boxed, positions of splice sites used to generate the various isoforms are indicated by vertical lines. B.) Predicted amino-acid sequences of the six Oct2 proteins. The full sequence of the major Oct2 form (Oct2.1) is shown, for the other isoforms only the divergent sequences are shown. Identical amino-acids are indicated by dashes (-), stop codons are indicated by asterisks. The POU-specific and the POU-homeo domain are boxed again, the 4 leucine residues that constitute the presumptive leucine zipper are hallmarkd by dots.
sequence of the human cDNA or the 3' end of the cloned murine cDNA to amplify Oct2 cDNA sequences by PCR. Surprisingly, we obtained alternate forms of both the 5' as well as the 3' end (data not shown). In order to determine which 5' ends would be associated with which 3' ends, we used the PCR technique to amplify the complete murine homologue of the published human cDNA (for experimental details, see Materials and Methods section). The amplified DNA was subcloned into a plasmid vector and 90 independent clones containing Oct2 sequences were selected for further analyses. Using frequently cutting restriction enzymes, we were able to classify about 80% of these cDNAs into 6 different forms. At least two isolates for each form were obtained. The remaining 20% of the cDNA clones were single isolates and not characterized further. More than half (63%) of the repeatedly isolated clones belonged to the class designated Oct2.1. The other forms were present at lower frequencies comprising between 3% (Oct2.6) and 12% (Oct2.3) of the clones. Figure 1 shows the composite nucleotide sequences and a summary of the deduced amino acid sequences of the 6 forms (Oct2.1–Oct2.6) characterized in this report. The dominant form (Oct2.1) is 98.5% identical to the human Oct2 clone isolated from Namalwa cells (29). Oct2.2 shares the same degree of identity with the Oct2 clones isolated from the human BJA-B cell line (28, 38). The difference between the two clones is due to an in frame insert of 48bp (16 aa) that are present in form Oct2.2 but not in Oct2.1 (nor in any of the other Oct2 variants). The other Oct2 variants have not yet been described. Oct2.3 has an in frame insert of 66bp (22aa) close to the amino terminus of the gene. Oct2.6 shows an in frame deletion of 117bp (39aa) starting at the insertion site of the extra 66bp of variant Oct2.3 and extending towards the carboxy terminus of the protein. The alterations in Oct2.4 and Oct2.5 are both localized 3' of the conserved POU-domain and change the reading frame. Oct2.4 lacks 136bp and runs into a stop codon 17aa downstream of the deletion site. This modification destroys the putative leucine zipper (28, 29, 38). Oct2.5 contains a 74bp insert 12aa upstream of the termination codon of variants Oct2.1, Oct2.2, Oct2.3 and Oct2.6. Due to this insertion and the resulting frame shift, this variant encodes 132 new amino acids at the carboxy terminus. The predicted molecular weights of the proteins encoded by the variant forms indicate that Oct2.1, Oct2.2 and Oct2.3 with molecular weights of approximately 57, 59 and 60kD could be considered Oct2A type proteins (21). With a predicted molecular weight of 75kD, Oct2.5 is a prime candidate to encode the previously described Oct2B protein (22). No lymphoid specific isoforms were detected (27, 28, 38); data not shown). At present, it is unclear whether the missing extensions are attached to the 5' or 3' end of the cloned cDNAs.

A comparison of the exon sequences of the genomic and cDNA clones revealed that the two in frame insertions in clones Oct2.2 and Oct2.3 as well as Oct2.4 are generated by using alternative splice acceptor sites (Figure 2B). Oct2.6 is generated by skipping exon 4. The 74bp insert present in the Oct2.5 isoform represents an autonomous miniexon that is spliced out in all the other isoforms (Figure 2B).

Both the POU-specific as well as the POU-homeo domains are split by an intron (Figures 1B and 2B). The position of the intron in the POU-specific subdomain does not correspond to the short stretch of amino acids that separates the POU-A and the POU-B regions (35) but rather splits the POU-A region. In addition, the leucine zipper region localized 3' to the POU-domain is also split by an intron (Fig. 2B).

The frequencies with which the different isoforms occurred in the pool of cloned cDNAs indicated that they might not be present in equimolar amounts in WEHI231 cells. Therefore RNase protection assays that allowed us to distinguish between the different Oct2 forms were performed to measure their relative frequencies (Fig. 3A). We were readily able to detect the appropriate protected fragments for isoforms Oct2.2, Oct2.3 and Oct2.5 as well as Oct2.4 in addition to Oct2.1 which represents the dominant form in WEHI231 RNA (Fig. 3B and 3C). The
Figure 3: RNA corresponding to the different isoforms are present at varying levels in B-cells. A.) Schematic representation of the fragments that are predicted to be protected from RNase degradation. The large fragment represents the Oct2 cDNA fragment subcloned into the pBluescript SK− vector used to generate the probe. The actual probe size is 50 to 60 nt longer than the size of the largest protected fragment due to the presence of polylinker sequences. The open boxes indicate the extra sequences present in the Oct2.2, Oct2.3 and Oct2.5 forms, respectively, the black box indicates the region missing in the Oct2.4 cDNA. Correspondingly for Oct2.2, Oct2.3 and Oct2.5 the largest protected fragment is indicative for the respective RNA whereas in the case of Oct2.4 the presence of the shorter protected fragment is indicative for the respective RNA. The other protected fragments are indicative for all the Oct2 RNAs except the specific form. B.) RNase protection with the Oct2.2, Oct2.3 and Oct2.5 probes using 30 μg cytoplasmic WEHI231 RNA. The marker shown in the left lane is an endlabelled MspI digest of pBR322 DNA. In all cases an additional fragment corresponding to residual undigested probe is seen above the largest protected fragment. C.) RNase protection with the Oct2.4 probe and RNA isolated from either WEHI231 cells, total spleen (50 μg each) or NIH/3T3 cells stably transformed with an Oct2.4 expression construct (20 μg). The positions of the correct protected fragments are indicated by arrowheads. As the stably transformed cells only express Oct2.4 the large protected fragment indicative for the forms Oct2.1–3 and Oct2.5–6 is absent in this lane.

two larger fragments in the Oct2.5 specific protection experiment show a somewhat aberrant running behavior by migrating too slowly in the gel system used (even more pronounced in Figure 6, see below). These bands represent the correct protected fragments, however, as comigrating bands were specifically obtained by RNase protection using RNA from fibroblast cells transfected with either Oct2.1 or Oct2.5 cDNA expression constructs (data not shown, for structure of the expression constructs see Figure 5A). RNA corresponding to isoform Oct2.6 could only be detected using a PCR protocol and an amplification primer specific for the indicative junction of the Oct2.6 cDNA (data not shown).

From densitometric scanning of the autoradiographs in Figure 3B and 3C we calculate that Oct2.2 is present at a frequency of roughly 5% of total Oct2 RNA whereas Oct2.3 and Oct2.5 make up about 11% and 18% of the Oct2 RNA, respectively. Oct2.4 represents about 8% and Oct2.6 is very infrequent making up less than 1% of the Oct2 RNA. From these numbers we estimate that Oct2.1 makes up about 55% and therefore represents the majority of the Oct2 RNA present in WEHI231 cells. These numbers are good reflections of the frequencies with which the clones were originally detected in the PCR cloning protocol (with the noted exception of Oct2.6) arguing for the general validity of the approach.

Oct2 isoforms are functional transcription factors

The deduced amino acid sequences shown in Figure 1 suggest that the different isoforms encode proteins of quite distinct sizes. This was verified both by overexpression of the individual isoforms in a bacterial expression system as well as by in vitro translation in reticulocyte lysates (data not shown). When used in an electrophoretic mobility shift assay (EMSA) specific complexes with an octamer containing probe were obtained with all six isoforms (Fig. 4). A comparison of the retarded complexes
Figure 4: *In vitro* synthesized Oct2 proteins specifically bind to the octamer motif. Electrophoretic mobility shift assays with 2μl aliquots of the *in vitro* translation reactions (lanes 1–14) or a BJA-B nuclear extract (lanes 15,16). A radiolabelled probe containing either a wild type (odd numbered lanes) or mutant octamer motif (even numbered lanes) was used. *In vitro* translation without added RNA (lanes 1,2), Oct2.1 (lanes 3,4), Oct2.2 (lanes 5,6), Oct2.3 (lanes 7,8), Oct2.4 (lanes 9,10), Oct2.5 (lanes 11,12) or Oct2.6 (lanes 13,14) RNA, respectively. The positions of the known Oct2 shifts (Octl, Oct2B and Oct2A) and the free probe (F) are marked. The complex migrating ahead of the Oct2A shift in the lanes with the BJA-B nuclear extract is most likely a non-specific complex, because it is also detectable with the mutant octamer probe.

with those obtained with a control BJA-B nuclear extract (Fig. 4, lanes 15 and 16) indicates that the proteins encoded by Oct2.1, Oct2.2 and Oct2.3 all shift the wild-type probe to a position similar to the Oct2A complex present in BJA-B nuclear extracts (Fig. 4, lanes 3,5,7 and 15). This indicates that the previous designation Oct2A does not describe a single protein but rather a family of similar sized proteins. The Oct2.5 protein shifts the probe to a position that corresponds to the position of the Oct2B band shift (compare lanes 11 and 15 in Fig. 4B). This comigration is still observed when the gels are run much further to yield a higher resolution (data not shown). This comigration in the EMSA together with the observed size of Oct2.5 in the SDS-polyacrylamide gel strongly suggests that Oct2.5 encodes the previously described Oct2B protein. Oct2.4 and Oct2.6 protein-DNA complexes migrate faster than the Oct2A protein-DNA complexes and have not been described previously (Fig. 4B, lanes 9 and 13). There is a complex of similar mobility visible with the BJA-B extract preparations, however (E. Schreiber, pers. communication). This complex might be hidden underneath the non-specific shift present in our extracts.

The ability of the different Oct2 isoforms to activate transcription was tested in a cotransfection assay. The human Oct2A cDNA (corresponding to our Oct2.2 isoform) had previously been shown to be able to stimulate an octamer promoter construct in non-lymphoid cells (28). Oct2 cDNAs were introduced into a cytomegalovirus enhancer/promoter based
eukaryotic expression vector. The AUG codon was provided by the vector and 6 amino acids encoded by the vector polylinker replaced the first two amino acids of Oct2 (Fig. 5A). The reporter plasmids containing either a wild-type or a point mutant version of the octamer motif upstream of the β-globin TATA-box and the modified β-globin gene have been described previously (10). The reporter plasmids also contain a modified mouse histone H4 gene that allows correction for different transfection efficiencies. NIH/3T3 fibroblast cells were cotransfected and RNA was isolated for quantitative analysis 48hrs later. From the results obtained it is evident that all 6 isoforms can stimulate the wild-type octamer promoter construct whereas the constructs containing a mutant octamer motif remain unaffected (Fig. 5B, lanes 1—12). Furthermore, it is obvious that the stimulation depends on the expression of the Oct cDNA because a control cotransfection with the expression vector lacking an Oct2 cDNA insert does not give this stimulatory effect (Fig. 5B, lanes 13,14).

Oct2 splicing pattern is conserved in murine B-cell development and also between species

The various Oct2 isotypes had all been isolated from the WEHI231 cell line which represents a surface immunoglobulin positive B-cell. It was of interest to determine whether B-cells at other differentiation stages would express the same set of Oct2 isoforms or whether there would be developmental stage specific alternative splicing of Oct2 mRNA. As representatives of different stages of the B-cell lineage we chose the HAFTL 1, 70Z/3 (early and late pre B-cells, resp.), WEHI231 (mature B-cells) and X63Ag8/653 (plasma cells) cell lines. RNase protection analysis of cytoplasmic RNA from the different cell lines with probes detecting Oct2.2, Oct2.3 and Oct2.5 in addition to Oct2.1 clearly shows that the Oct2 splicing pattern is conserved throughout the B-cell lineage (Figure 6A). Furthermore it is evident that although the overall quantities of Oct2 RNA are lower in the cell lines representing the earlier stages of B-cell development, the relative frequencies of the individual isoforms are roughly the same in all cell lines tested. The same result was obtained when a probe specific for Oct2.4 was used or PCR analysis with an Oct2.6 specific primer was performed (data not shown).

We also analyzed Oct2 expression in different lymphoid organs to exclude the possibility that the observed alternative splicing is the consequence of continuous growth of cells in culture. RNase protection analysis of total RNA isolated from bone marrow, thymus, spleen, and mitogen stimulated spleen cells reveals that in all these primary samples the multitude of Oct2 isoform expression is conserved (Figure 6B). Again the relative frequencies of different Oct2 isoforms are comparable whereas the overall amounts of Oct2 RNA vary in the different organs. The detection of the full range of Oct2 isoforms in bone marrow RNA is consistent with the previous observation that pro and pre B-cell lines already express the various Oct2 isoforms. The presence of Oct2 RNA in thymus and ConA stimulated spleen cells suggests that primary T-cells actually express Oct2. Clearly the strongest expression of Oct2 is observed in spleen and lipopolysaccharide (LPS) stimulated spleen cells. However, it is interesting that mitogen stimulation does not lead to a significant increase of Oct2 RNA expression.

The previous isolation of two Oct2A cDNAs from the human BJAB cell line corresponding to the mouse Oct2.1 and Oct2.2 isoforms indicated that the alternative splicing pattern is conserved even between mouse and human. To further extend this observation we investigated whether the mouse Oct2.5/Oct2B

Figure 6: Expression of Oct2 isoforms in different B-cell lines and lymphoid organs. A.) RNase protection analysis with the Oct2.2 (lanes 2—5), Oct2.3 (lanes 7—10) and Oct2.5 (lanes 12—15) specific probes. 30μg cytoplasmic RNA from HAFTL1 (lanes 2,7,12), 70Z/3 (lanes 3,8,13), WEHI231 (lanes 4,9,14) and X63Ag8/653 cells (lanes 5,10,15) were used. The position of the correct protected fragments is indicated by arrowheads. The marker lanes are endlabelled Mspl digests of pBR322 DNA. B.) RNase protection analysis with RNA from various organs. The same probes as above were used (Oct2.2: lanes 2—7; Oct2.3: lanes 8—13; Oct2.5: lanes 14—19). 25μg of cytoplasmic WEHI231 RNA (lanes 2,8,14) or 50μg total RNA from the following sources were used: bone marrow: lanes 3,9,15; thymus: 4,10,16; spleen: lanes 5,11,17; spleen/LPS: lanes 6,12,18; spleen/ConA: lanes 7,13,19;
The octamer motif turns out to be a multi-functional regulatory element recognized by a variety of different proteins. So far, at least a subset of T-lymphocytes.

Multiplicity of octamer binding proteins
The octamer motif turns out to be a multi-functional regulatory element recognized by a variety of different proteins. So far, at least three genes encoding octamer binding proteins (Oct1, Oct2 and Oct4) have been cloned (17, 24, 27, 28, 29, 30, 31). Additional octamer binding proteins have been identified, but it is at present unclear whether they are encoded by separate genes (23). Multiple genes coding for distinct transcription factors interacting with a conserved motif have been described for proteins binding to the CCAAT-box, the AP-1 site, and the cAMP response element (CRE) (44, 45, 46, 47, 48, 49). We show here that alternative splicing is responsible for the generation of a multitude of Oct2 isoforms. Preliminary evidence suggests that the Oct4 gene might also code for two protein isoforms, Oct4 and Oct5, due to the usage of alternate translational initiation codons (30). Alternative splicing as a mechanism to generate multiple isoforms of transcriptional regulators has also been described for the CTF/NF-1 transcription factor (45), the CREB transcription factor (50), as well as for the drosophila ultrabithorax (Ubx) gene (51). Differential activities for alternate forms have been described for both CREB and CTF/NF-1 (50, 52). Another striking homology between the CCAAT-box and octamer binding proteins is the fact that members of either family can function in stimulation of both transcription and replication (52, 53).

DISCUSSION
We have shown that the lymphoid-specific Oct2 transcription factor family is expressed as multiple alternatively spliced isoforms in mouse and human B-cells. All the isoforms described in this paper retain the conserved POU-domain and are therefore able to bind specifically to the cognate octamer motif. Furthermore all the splicing variants retain the ability to activate an octamer containing promoter in a cotransfection assay. Our analyses also show that the pattern of alternative splicing is conserved throughout the murine B-cell lineage and probably also in at least a subset of T-lymphocytes.

Possible functions of the various Oct2 isoforms
The Oct2 transcription factors could be differentially involved in several aspects during B-cell ontogeny. One of the first characteristics of a committed precursor of the B-cell lineage is the transcription of unrearranged heavy-chain genes (sterile μ transcription). It has recently been demonstrated that the octamer element in the heavy-chain enhancer plays an important role for this sterile μ transcription (54). The next important step in B-cell development is the rearrangement of the heavy-chain locus. Binding of Oct2 to its recognition sites in the heavy-chain enhancer and the V_{H}-promoters could be a critical event preceding this rearrangement. In this respect it is of interest that the V_{H}-promoters contain a high- and a low-affinity binding site for Oct2, the octamer and the heptamer motif (55, 56), and Oct2 binding to these two sites shows cooperativity (43, 57). Although this cooperative binding has been shown in vitro using purified bacterially expressed Oct2 protein corresponding to Oct2.2 (43) it is not known what type of Oct2 protein binds to these motifs under in vivo conditions.

If different Oct2 isoforms have different functions during B-cell development one might have expected to observe changes in the Oct2 RNA splicing pattern. Although our analyses clearly show that this is not the case, this does not exclude functional differences of the various isoforms. One possibility could be that the Oct2 isoforms are regulated specifically at the translational level. Interestingly, expression of the GFH-1/Pit-1 protein, another member of the POU domain gene family, has been suggested to be regulated at the translational level in development (58). Differential activation properties could also be the result of differential modifications of the various isoforms. It has recently been reported that differential phosphorylation of Oct1 and Oct2A are due to differences at the carboxy termini of the two proteins and that these differences correlate with the transcriptional potential (59). Members of the Oct2 transcription factor family might also be recruited for specific functions as a consequence of specific protein-protein interactions. Interestingly, isoform Oct2.4 is devoid of a functional leucine zipper, a structure that has been shown to be involved in protein-protein interactions (60, 61, 62, 63). This model has been suggested to explain the differential activities of the Oct1 and Oct2 proteins which are coexpressed in lymphoid cells (64).

The observation that all the various Oct2 isoforms stimulate an octamer containing promoter in a cotransfection assay shows
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