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# **Open Access** Cloning and functional characterisation of avian transcription factor E<sub>2</sub>A Thomas M Conlon and Kerstin B Meyer\*

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# Abstract

Background: During B lymphocyte development the E2A gene is a critical regulator of cell proliferation and differentiation. With regards to the immunoglobulin genes the E2A proteins contribute to the regulation of gene rearrangement, expression and class switch recombination. We are now using the chicken cell line DT40 as a model system to further analyse the function of E2A.

**Results:** Here we report the cloning and functional analysis of the transcription factor E2A from chicken. Using RACE PCR on the chicken lymphoma cell line DT40 we have isolated full-length clones for the two E2A splice variants E12 and E47. Sequence conservation between the human and chicken proteins is extensive: the basic-helix-loop-helix DNA binding domain of human and chicken E47 and E12 are 93% and 92% identical, respectively. In addition high levels of conservation are seen in activation domain I, the potential NLS and the ubiquitin ligase interaction domain. E2A is expressed in a variety of tissues in chicken, with higher levels of expression in organs rich in immune cells. We demonstrate that chicken E12 and E47 proteins are strong transcriptional activators whose function depends on the presence of activation domain I. As in mammals, the dominant negative proteins Id1 and Id3 can inhibit the function of chicken E47.

Conclusions: The potential for homologous recombination in DT40 allows the genetic dissection of biochemical pathways in somatic cells. With the cloning of avian E2A and the recent description of an in vitro somatic hypermutation assay in this cell line, it should now be possible to dissect the potential role of E2A in the regulation of somatic hypermutation and gene conversion.

# **Background**

The transcription factor E2A contributes to transcriptional regulation in many cell lineages. However, it is essential for the development of B lymphocytes [1,2]. Its role in mammalian B cell development has been studied extensively and E2A functions in B cell commitment and proliferation as well as immunoglobulin (Ig) gene rearrangement and expression (reviewed in [3]). By alternative splicing the E2A gene encodes two basic helix-loophelix (bHLH) proteins, E12 and E47, which differ only in their highly homologous DNA binding and dimerisation domains [4]. Binding sites (consensus CANNTG) for these transcription factors are found in all Ig enhancers as well as a number of genes required for heavy and light chain rearrangement ( $\lambda 5$ , rag-1, rag-2, EBF). Remarkably, over-expression of E2A together with the recombinase activating genes is sufficient to allow rearrangement of the endogenous Ig locus in a non-lymphoid cell line [5].

E2A also plays a role in peripheral B cell differentiation. E2A protein expression is greatest in the highly proliferative dark zones of germinal centres, where class switching (CSR) and somatic hypermutation (SHM) are thought to occur. In keeping with such a role, repression of E2A via Id proteins inhibits CSR [6] and leads to lower expression of the AID (activation induced deaminase) gene [7], a gene that is essential for both CSR and SHM to occur [8,9].

Avian B cell development differs from mammalian B cell development in a number of important aspects. Rearrangement occurs between a single V<sub>L</sub> and J<sub>L</sub> segment to yield a single functional variable light chain gene. Similarly a single  $\mathrm{V}_{\mathrm{H}}$  segment combines with one of 15 D and a single J segment [10] generating only a small repertoire. Gene conversion subsequently utilises a pool of upstream pseudo V-genes to generate diversity [10,11]. Thus gene conversion is the primary mechanism to establish the B cell repertoire in chickens. The diversification and expansion of B cell progenitors occurs in the specialised microenvironment of the Bursa of Fabricus from where mature B cells exit into the periphery. By 6–8 months the diversity of the B cell compartment is established and the bursa involutes. Given the distinct nature of avian B cell development we were interested to find out whether the transcription factor E2A plays a similar central role in B cell ontogeny and the generation of diversity.

To this end we have now cloned full-length avian E12 and E47 cDNAs from the chicken B cell lymphoma DT40 and have initiated their functional characterisation.

# Results and Discussion Cloning of chicken E12 and E47

A chicken bursal EST database [12,13], was searched with the bHLH domain of human E47, revealing a single highly homologous clone of 785 bp. Using a combination of degenerate and RACE PCR a 2.47 kb sequence encoding full-length chicken E47 was obtained. The protein sequence downstream of the first methionine is highly conserved across human, mouse, xenopus and chicken (Fig. 1A) suggesting that the true 5' end has been identified. Furthermore, nucleotides 3 bp and 6 bp upstream of the ATG correspond to the Kozak consensus sequence [14]. In order to address whether chicken, like its mammalian homologues, contains the alternatively spliced E12 DNA binding exon, primers were designed on either side of the bHLH region (P7, 8) and used in RT-PCR on DT40 cDNA. The resultant clones contained either E12 (542-620aa) or E47 (542-616aa) sequence (boxed sequences in Fig. 1A and 1B), while the sequence flanking the bHLH exon was identical.

# Sequence analysis

The resultant full-length E12 and E47 protein sequences were compared to the human, mouse and xenopus orthologues using CLUSTAL W [15], and the results of this analysis are depicted in Fig. 1. Chicken E12 and E47 are identical except for their bHLH domain, which is nevertheless highly homologous with 72% identity and 86% similarity when allowing only conservative changes. Compared to their human orthologues, chicken E47 and E12 are 71% and 69% identical, respectively. Within the bHLH domains the conservation is even greater. Chicken and human E47 bHLH are 93% identical and chicken and human E12 bHLH are 92% identical. Thus E47 bHLH domains of different species are more similar than the E12 and E47 domain within one species. With regard to the other species examined, chicken E12 is slightly more homologous to xenopus E12 (70%) than to mouse E12 (68%). Outside the bHLH domain there are a number of well-conserved domains. In particular, activation domain I (ADI) is highly homologous across all four species examined (see Fig. 1). The chicken sequence corresponds to the helix consensus described for interaction with the SAGA complex [16], indicating that E2A mediates some of its function at the chromatin level. Activation domain II has been identified as a domain required to drive transcription in insulin producing  $\beta$ -cells [17]. Within activation domain II only the 5' region is conserved, while insertions are found within the 3' region. However the length of the conserved region, which is significantly longer than that of the ADI, may indicate that this domain has additional functions. The putative nuclear localisation domain is identical in all the compared sequences. Another functional region of the E47 protein is the domain interacting with ubiquitin conjugating enzyme UbcE2A (477-530aa [18,19]). Within this region, runs of very high homology are seen between the sequences. Whether this is sufficient to obtain functional interaction with UbcE2A awaits experimental confirmation.

In conclusion all regions of very high sequence conservation correspond to functionally important domains.

# Genomic structure of the E2A gene

We have mapped the intron-exon structure of the E2A gene on the most recent release of the chicken genome using the genome browser at UC Santa Cruz [20]. Chicken E2A is encoded on chromosome 28 and comprises 19 exons spanning 38.7 kb. A schematic representation of the gene is shown in Fig. 2. The intron-exon boundaries have been mapped (Table 2, see Additional file 1) and all introns conform to the splice consensus sequence. Three exons (2, 5 and 7) lie in gap sequences of the genome and searches with these exons did not reveal any homologies in the chicken genome. Their exact location has therefore not been assigned. However, the length of these three

A	Alignment of E12	human mouse xenopus <b>chicken</b>	-MNQP-QRMAPVGTDKELSDLLDFSMMFFLPVTNGKGRPASLAGAQFGG-SGLEDRPSSG MMNQS-QRMAPVGSDKELSDLLDFSMMFFLPVANGKSRPASLGGTQFAG-SGLEDRPSSG -MNQ-QQRMTAVGTDKELSDLLDFSMMFFLPVANGKNRPTTLASTQFGG-SGLEERTGSA -MNQQQGNMAAVGTDKELSDLLDFSMMFFLPVANGKNRPTTLASTQFGG-SGLEERTGSG	57 58 58 58
			*** ***:.**:***************************	
			helix of ADI	
		human	SWGSGDQSSSSFDPSRTFSEGTHFTESHSSLSSSTFLGPGLGGKSGERGAYASFG	112
		mouse	SWGSSDQNSSSFDPSRTYSEGAHFSDSHSSLPPSTFLGAGLGGKGSERNAYATFG PWGTEDHNGSTFDOGRGYGDGTHYGEHRELPSHEGNLSAPELASGLVGKN-DRPPYSTFG	113
		chicken	SWGTGDQNSSTFDQGRSYGEGPHYGEHRDLPSHNSLSSSPFLGPGLVGKSGERASYSTFG	118
			.**: *:*:** .* :.:*.*: : *:.**** **. :* .*::**	
		human	RDAGVGGLTQAGFLSGELALNSPGPLSPSGMKGTSQYYPSYSGSSRRRAADGSLDTQPKK	172
		mouse	RDTSVGTLSQAGFLPGELSLSSPGPLSPSGIKSSSQYYPSFPSNPRRRAADGGLDTQPKK	173
		xenopus	RDSGMT-MNQTGFLSGDLAMNSPSALSPNAGKAGSQYY-TYPNNSRRPGDSNLDAQPKK	175
		Chicken	**:.: :.*:***::: *****. *. ***: ::****	1//
		human	VRKVPPGLPSSVYPPSSGEDYGRDATAYPSAKTPSSTYPAPFYVADGSLHPSAELWSPP-	231
		mouse	VRKVPPGLPSSVYPPSSGDSYSRDAAAYPSAKTPSSAYPSPFYVADGSLHPSAELWSTP-	232
		xenopus	VRKVPPGLPSSVYPANSGDEYCGDRGSYASTKTPNSVYPGAFYMADG-LHSS-DIWASSG	233
		chicken	VRKVPPGLPSSVYPSNSGDDYSRDPAGYTPSKPPSTVYPGAFYMADG-LHNSPDLWSSPG	236
		human	GOAGEGPMLGGGSSPLPLPGSGPVGSSGSSSTEGGLHOHERMGYOLHGAEVNGGLPS	289
		mouse	SQVGFGPMLGDGSSPLPLAPGSSSVGSGTFGGLQQQDRMGYQLHGSEVNGSLPA	286
		xenopus	GIGQGGYSSVLNSSQSSVSQTTGFSSLHPHERMGFPMHSGEVNPS	278
		Chicken	* .::** ***.: **	284
		human	ASS-FSSAPGATYGGVSSHTPPVSGADSLLGSRGTTAGSSGDALGKALASIYSPDHSS	346
		mouse	VSS-FSAAPG-TYSGTSGHTPPVSGAAAESLLGTRGTTASSSGDALGKALASIYSPDHSS	344
		xenopus	VTSSFSSTPAQYGVSSHTPPISTGDTIIGNRGTATGSSGDALGKALASIYSPDHSS	334
		Chicken	**::. *.*.***:* ::::***************	555
		human	ADII NNFSSSPSTPVGSPQGLAGTSQWPRAGAPGALSPSYDGGLHGLQSKIEDHLDEAIHVLRS	406
		mouse	NNFSPSPSTPVGSPQGLPGTSQWPRAGAPSALSPNYDAGLHGLQSKMEDRLDEAIHVLRS	404
		xenopus chicken	TNFSSTPSTPVGSPQGITGSGQWPRANGPGALSPSYDGTLHTLQNKMEDRLDEAIHVLRS NNFSSNPSTPVGSPOGLAGTSOWSRAGGOGALSPSYEGSLHTLONKMEDRLDEAIHVLRN	394 399
			·***********************************	
		human	HAVGTAGDMHTLLPGHGALASGFTG-PMSLGGRHAGLVGGSHPEDG	451
		mouse	HAVGTASDLHGLLPGHGALTTSFTG-PMSLGGRHAGLVGGSHPEEG	449
		chicken	HAVGQIGALFGGADDINSLLSAIASVLGSGFFAAVLSLAWARSALM-SHIELA HAVGQIAAMPSNHGDMHGLLGSAPAHSAAVGSLGQAFPASVMSLGGRHPSLVGGGHPEDG	459
			**** :. *:* ** * .* :****. * .*.*:.	
		human	LAGSTSLMHNHAALPSQPGTLPDLS-RPPDSYSG-LGRAGATAAASEIKREEKEDEE	506
		mouse	LTSGASLLHNHASLPSQPSSLPDLSQRPPDSYSG-LGRAGTTAGASEIKREEKEDEE	505
		chicken	LSSTPSMLHNHVTLPSQPSSLPDLN-RQQDTYSGLSGGLGRSSVSSGTNEIKREEKEDEE	518
			*::*.:::*::*::*: * *:** * ***: :****: Ubc-L	
		human	NTSAADHSEEEKKELKAPRARTSPDEDEDDLLPPEQKAEREKERRVANNARE	558
		mouse	1ASVAD-AEEDKKDLKVPRTRTSPDEDEDDLLPPEQKAEREKERRVANNARE NRSVADLSDDEKKESKPORSRTRCSINSDDEDEDEDDDNLPPEOKAEREKERRVANNARE	556
		chicken	NTSVADNSEEEKKELKPSRNRTS <u>ODED</u> EEDDLLPPEQKAERERERRVANNARE	571
			*.** ::::**: * * ** * *****************	
		human	RLRVRDINEAFKELGRMCQLHLNSEKPQTKLLILHQAVSVILNLEQQVRERNLNPKAACL	618
		xenopus	RLRVKDINEAFKELGRMCQLHLNSEKPQTKLLILHQAVSVILSLEQQVPERNLDPKAACL	624
		chicken	RLRVRDINEAFKELGRMCOLHLNSEKPOTKLLILHOAVSVILNLEOOVF	631
		human	KRREEEKVSGVVGDPQMVLSAPHPGLSEAHNPAGHM 654	
		mouse	KRREEEKVSGVVGDPQLALSAAHPGLGEAHNPAGHL 652	
		xenopus chicken	KRREEEKVSGVDPQMGLSGGHPGVGDSHNPVGHM 658 KRREEEKVSGVUGDPOMTLSASHPGLGDGHNPVGHM 667	
		cm chen	********** ***: ***: ***::::***:**:	
Р	۸. ا: مربع مربع مربع ا	human 530	STDEVLSLEEKDLRDRERRMANNARERVRVRDINEAFRELGRMCQMHLKSDKA	582
В	Alignment	mouse 528	STDEVLSLEEKDLRDRERRMANNARERVRVRDINEAFRELGRMCQLHLKSDKA	580 591
	of F47	chicken 542	STDEALSLEDKDLRDRERRMANNARERVRVRDINEAFKELGRMCQMHLKTDKA	594
			** . :*:** :*****:*********************	
	DHLH exon	human	QTKLLILQQAVQVILGLEQQVR 604	
		mouse	QTKLLILQQAVQVILGLEQQVR 602	
		chicken	QTKLIILQQAVQVILGLEQQVR 616	
			**** **** ** ** *****	

#### Figure I

Alignment of E2A sequences from human, mouse, xenopus and chicken: full-length E12 (**A**) and the E47 bHLH exon (**B**) which replaces the E12 bHLH exon in the full-length sequence. "\*" denotes identical residues, ":" conserved substitutions as recorded in CLUSTAL, and "." semi-conservative substitutions. Accession numbers for the sequences used are: PI5923, XM125750, X66959 for human, mouse and Xenopus, respectively. The chicken sequence has been assigned the accession number AJ579995 for E12 and AJ579996 for E47. Shaded boxes denote functionally defined regions: activation domain I (ADI) with boxed helix [29], activation domain II (ADII) [17] and the ubiquitin ligase interaction domain [18]. The putative NLS is underlined and the bHLH region is boxed in black. The exon border of the xenopus sequence is based on its homology to the other sequences. Chicken amino acid 221 (bold) is the first aa of the  $\Delta$ ADI constructs used in the functional analysis.



#### Figure 2

Schematic representation of the structure of the chicken and human E2A genes. Black lines represent assigned exons, grey lines represent exons lying in gap sequences; their exact position has not been determined. Open boxes represent the alternatively spliced E12 and E47 exons encoding the helix-loop-helix domain of these proteins. See Table 2 for the exact position of each of the exons.

exons is very similar to exons 2, 5 and 7 in the human gene (see Table 2) suggesting that they are indeed encoded as single exons. The overall structure of the human and chicken genes is extremely well conserved, with the human gene containing 18 exons spanning 40.98 kb of chromosome 19 (Fig. 2). The intron-exon borders are very similar, the only exception being the 3'UTR which is split into 2 exons in the chicken whilst the 3' UTR is encoded as a single exon in the human gene. In both chicken and human the first 2 exons are widely spaced, with exons 3 lying 21.5 kb upstream of the ATG in the chicken and 17.9 kb in the human E2A gene. In contrast the exons at the 3' end of the gene are highly clustered with exons 8–19 spanning 9.2 kb in the chicken and 10.4 kb in the human gene.

Furthermore we found three clusters of highly related sequence between the chicken and human loci that did not correspond to coding sequence. These mapped to upstream and downstream of the gene and to a region upstream of exon 3. We are currently investigating whether these function as regulatory sequences.

#### Chicken E2A expression pattern

To examine whether there is differential expression between E12 and E47 in the chicken DT40 cell line, splice variant specific primers (P10, P11 and P12) were used in semi-quantative RT-PCR of DT40 mRNA. Fig. 3A demonstrates that the relative amounts of E12 and E47 expressed are very similar. Furthermore we examined E2A gene



# Figure 3

**A.** PCR analysis of E12 and E47 on DT40 cDNA using increasing dilutions of cDNA. M, DNA markers. **B.** Northern Blot of E2A expression, with 2  $\mu$ g polyA+ RNA from the tissues shown above the lanes. A  $\beta$ -actin probe was used as a control.

expression in different chicken parenchymal tissues. As observed in human and mouse, low levels of E2A expression are detected by Northern blotting in many different tissues in the chicken (Fig. 3B). Of the tissues examined, greatest expression is detected in the lung, spleen and ileum, sites where immune cells accumulate. However, relatively high expression is also seen in testis and the ovaries. The size of the E2A message is approximately 2.5 kb which corresponds well to the length of cDNA clones we have isolated. However, in spleen and to a lesser extent in ileum and testis there is a second smaller message of about 2.2 kb. The functional significance of this is not yet clear. The overall expression pattern is in keeping with E2A contributing to gene expression in a variety of tissues but being of particular importance for lymphocytes. Like in mammalian cells the wide expression pattern may not reflect protein levels since E2A is regulated post-transcriptionally [6,21].

# Functional analysis of chicken E2A

Having cloned chicken E2A we wished to confirm its ability to activate transcription from E2A containing reporter plasmids. We demonstrate that chicken E47 and E12 strongly activate transcription from a multimerised E2A binding site linked to a TATA box (Fig. 4A). Upon transfection into human HEK293 cells, hamster E47 (MDE47 [22]) stimulated transcription by 1000-fold. Chicken E12 and E47 gave similar values with E12 being approximately 20% less active than E47. For both isoforms transcriptional activation strongly depended on the presence of its 5' end, including activation domain I (ADI). E12ΔADI retains less than 1% of the activity of the full-length protein, while E47 $\Delta$ ADI is reduced to 3.5% of its wild type activity. The deleted region encompasses the ADI and the putative NLS. However, previous studies have shown that similar deletions of human E2A are nevertheless found in the nucleus [17]. This suggests that the loss of function in the mutant proteins is due to the deletion of activation domain I. Furthermore we investigated whether Id proteins, dominant negative regulators of bHLH function, can regulate chicken E2A proteins. To this end we amplified mouse Id1 [23] and mouse Id3 [24] by PCR and subcloned the resultant fragments into an expression vector, driven by the cytomegalo virus promoter/enhancer region, and tested their ability to abrogate transcriptional activation (Fig. 4B). Once again E2A transactivation was assayed on a multimerised E2A binding site. Under the experimental conditions used this yielded 4500-fold activation by E47. As before E12 activity was about 80% of that seen for E47. Co-transfection of mouse Id1 reduces E47 driven transcription to 10% of that seen for the control vector only, while the mouse Id3 was able to reduce transcription to 26%. Thus, like their mammalian orthologues chicken E2A proteins can be regulated by the Id proteins. Orthologues of the Id proteins in chicken have been cloned and their embryonal expression pattern examined [25]. As yet little is known about their expression in avian lymphoid development. However, our results suggest that Id proteins modulate the activity of chicken E2A proteins.

# Conclusions

Here we report the cloning of the chicken orthologue of the human transcription factor E2A. The identification of chicken orthologues of genes known to be important for the development of the immune system is of great interest with regard to understanding a related but distinct immune system as is found in birds. One of the advantages of utilising chicken as an experimental system is the existence of the bursal cell line DT40, which undergoes homologous recombination with high frequency thus allowing a genetic approach to the dissection of biochemical pathways. We plan to utilise DT40 in the analysis of E2A function in mature B cells. In mammalian cells, E2A is known to contribute to the regulation of class switch recombination and it has been suggested that it may contribute to SHM either directly or indirectly through controlling AID gene expression. Normally IgV gene diversification in chicken occurs by gene conversion,



### Figure 4

Transcriptional activation by mammalian E47 and chicken E12 and E47 in HEK293 cells. **A.** Transactivation of a multmerised E2A binding site by different E2A proteins as indicated. Values are given relative to those obtained with the pcDNA control vector. The data shows the mean +/- SEM of triplicate transfections. **B.** Effect of co-transfection of Id expression constructs on E47 transcriptional activation. The data show the mean +/- SEM of five independent transfections.

Table	e I:

Primer name	Sequence, 5' to 3'
PI	CTGGCCCACGGCGTGATTCCTCAAG
P2	GAAAGTTCCGGAGTGGTGGGAGACC
P3	GGGGTTTACTGCGAAAAGTCGAAG
P4	AGTGACCTCCTGGACTTCAG
P5	GGATCCCTGCTGTAGTCATCACCTGAG
P6	CTCCAAACTGAGTGCTTGCTAGGGTCG
P7	CAGGGATCCGGCTGGATAC
P8	AAGCTTAGGATCTTTACATGTGTCC
P9	GGTACCCAGGACTGCCACTGTTGAG
PI0	CTACCAGCTGCATTCAGGAGAG
PH	TCCTCCTCATCCTCATCTTG
P12	GATAAGGCCTCGTCTGTACTG
PI3	ATGAAGGTCGCCAGTGGCAG
PI4	TCAGCGACACAAGATGCG
P15	GTGCTGTGTTCCCATCTATCGT
PI6	TGGACAATGGAGGGTCCGGATT
P17	ATGAAAGTCGCCAGTGGCAG

Sequence of primers used in PCR reactions.

although somatic mutation can occur [26]. However, cells carrying mutations in the Rad51 homologues XRCC2 and XRCC3 undergo SHM with very high frequency [27]. The identification of avian E2A, together with the recent description of an in vitro somatic mutation assay in DT40 cells, should allow us to investigate the exact role E2A plays in regulating the complex events that confer specificity on the SHM mechanism.

# Methods

### RACE PCR and sequencing

Primers P1, P2 and P3 (for all primers see Table 1) were used in 5' and 3' RACE PCR as described in the SMART RACE cDNA Amplification Kit (BD Biosciences, USA) on DT40 total RNA (RNeasy, Qiagen, UK). Further PCR reaction with primers P4, P5 and P6 generated full-length E47 cDNA. All PCR products were cloned into the pCR2.1-TOPO vector (Invitrogen, USA) and sequenced using the ABI Big Dye terminator cycle sequencing ready reaction kit (Applied Biosystems, UK). Sequencing reactions were analysed in an ABI 3700 DNA Analyser (Applied Biosystems, UK).

# E2A expression

PCR reactions on serial dilutions of DT40 cDNA prepared using the SMART RACE cDNA Amplification Kit, were carried out using primer pairs P10 and P11 for E12 expression and P10 and P12 for E47 expression. 2  $\mu$ l of cDNA was used in 20  $\mu$ l reaction mixtures containing 0.2 mM of each dNTP, 2  $\mu$ l 10x PCR buffer (Applied Biosystems, UK), 1.5 mM MgCl<sub>2</sub> (E12) or 2 mM MgCl<sub>2</sub>(E47), 1  $\mu$ M of each primer and 0.5 U Taq Gold (Applied Biosystems, UK). PCR conditions were 94°C for 10 min followed by 35 cycles of 94°C for 30 s, 55°C for 30 s and 72°C for 3 min, followed by a 10 min extension at 72°C. Products were analysed on a 1.5% agarose gel. A NeverFail Northern Blot (RNway Laboratories) of 2 µg chicken poly A+ RNA per lane was probed with a <sup>32</sup>P-labelled fragment of E2A genomic DNA, overlapping the two exons from 1454–1576 and 1577–1726 of the cDNA. The β-actin control probe was generated by PCR using primers P15 and P16.

# Plasmids

Full length cDNAs for expression cloning of E12 and E47 were generated by a 2-step PCR reaction using primer pairs P7, P8 (3' end of proteins) and P9, P5 (extending to 5' start). Clones were confirmed by sequencing and cloned into the pcDNA3 expression vector (Invitrogen, USA) using BamH I and Not I to generate AAD1 E12 and E47 constructs. Both E12 and E47 are truncated at amino acic 221 (indicated in bold in figure 1). Subsequently the 5' ends were inserted using Kpn I and BamH I to generate full length clones. The MDE47 vector was obtained from M. Sigvardsson and contains a forced dimer of hamster E47 in pcDNA [22]. The 6xE2A luciferase vector (pXp2Luc) from A. Green's laboratory contains 3 copies of the following sequence: gtcgaacagatgttcacacgaccatctgtgg. pGL3-control and promoter vectors are from Promega, UK. Mouse Id1 was amplified using the primer pairs P13, P14 and cloned into the pCR2.1-TOPO vector. Id1 was subcloned into pcDNA3.1/Hygro expression vector (Invitrogen, USA) using BamH I and Not I. Full length mouse Id3 [28] was subcloned into the same expression vector using BamH I and HindIII.

# Cell culture and transfection

The chicken bursal B cell line DT40 cells was obtained from Dr J. Sale and maintained in RPMI-1640, 50  $\mu$ M  $\beta$ mercaptoethanol, 7% foetal calf serum, 3% chicken serum and antibiotics; human embryonic kidney cells, HEK293, were from Dr K.J. Patel and maintained in DMEM, 10% foetal calf serum and antibiotics. 1  $\mu$ g each of luciferase reporter and transactivation plasmids, and 0.25  $\mu$ g of CMV- $\beta$ -galactosidase vector, were transfected using Superfect Reagent (Qiagen, UK) according to the manufacturer's instructions. We harvested cells 36–48 hours after transfection and generated cell extracts in 200  $\mu$ l Promega reporter lysis buffer. Luciferase and  $\beta$ -galactosidase activity in 5  $\mu$ g of protein was measured using Promega reagents. Results are given as ratios of luciferase over  $\beta$ -galactosidase activity.

# **Authors' contributions**

TMC performed the cloning, sequencing and expression studies. KBM initiated the project, carried out some of the functional and the sequence analysis and drafted the manuscript. All authors have read and approved the final manuscript.

# **Additional material**

# Additional File 1

Table 2 Location of the chicken E2A exons, and exon and intron sizes of the chicken and the human E2A genes. Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2172-5-11-S1.doc]

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#### References

- Bain G, Maandag ECR, Izon DJ, Amsen D, Kruisbeek AM, Weintraub BC, Krop I, Schlissel MS, Feeney AJ, van Roon M, van der Valk M, te Rile JPJ, Berns A, Murre C: E2A proteins are required for proper B cell development and initiation of immunoglobulin gene rearrangements. *Cell* 1994, 79:885-892.
- Zhuang Y, Soriano P, Weintraub H: The helix-loop-helix gene E2A is required for B cell formation. Cell 1994, 79:875-884.
- Kee BL, Quong MW, Murre C: E2A proteins: essential regulators at multiple stages of B-cell development. *Immunol Rev* 2000, 175:138-149.
- Murre CP, McCaw PS, Baltimore D: A new DNA binding and dimerization motif in immunoglobulin enhancer binding, daughterless, myoD, and myc proteins. *Cell* 1989, 56:777-783.
- Romanov WJ, Langerak AW, Goebel P, Wolvers-Tettero ILM, van Dongen JJM, Feeney AJ, Murre C: E2A and EBF act in synergy with the V(D)J recombinase to generate a diverse immunoglobulin repertoire in nonlymphoid cells. *Mol Cell* 2000, 5:343-353.
- 6. Quong MW, Harris DP, Swain SL, Murre C: **E2A** activity is induced dring **B-cell** activation to promote immunoglobulin class switch recombination. *EMBO J* 1999, **18**:6307-6318.
- Sayegh CE, Quong MW, Agata Y, Murre C: E-proteins directly regulate expression of activation-induced deaminase in mature B cells. Nat Immunol 2003, 4:586-593.
- Muramatsu M, Kinoshita K, Fagarasan S, Yamada S, Shinkai Y, Honjo T: Class switch recombination and hypermutation require activation-induced cytidine deaminase (AID), a potential RNA editing enzyme. *Cell* 2000, 102:553-563.
- Revy P, Muto T, Levy Y, Geissmann F, Plebani A, Sanal O, Catalan N, Forveille M, Dufourcq-Labelouse R, Gennery A, Tezcan I, Ersoy F, Kayserili H, Ugazio AG, Brousse N, Muramatsu M, Notarangelo LD, Kinoshita K, Honjo T, Fischer A, Durandy A: Activation-induced cytidine deaminase (AID) deficiency causes the autosomal recessive form of the Hyper-IgM syndrome (HIGM2). Cell 2000, 102:565-575.
- Reynaud CA, Dahan A, Anquez V, Weill JC: Somatic hyperconversion diversifies the single Vh gene of the chicken with a high incidence in the D region. *Cell* 1989, 59:171-183.
- Reynaud CA, Anquez V, Grimal H, Weill JC: A hyperconversion mechanism generates the chicken light chain preimmune repertoire. Cell 1987, 48:369-378.
- Abdrakhmanov I, Lodygin D, Geroth P, Arakawa H, Law A, Plachy J, Korn B, Buerstedde JM: A large database of chicken bursal ESTs as a resource for the analysis of vertebrate gene function. *Genome Res* 2000, 10:2062-2069.
- 13. [http://swallow.gsf.de/dt40est.html]: .
- Kozak M: The scanning model for translation: an update. J Cell Biol 1989, 108:229-241.
- 15. [http://www.ebi.ac.uk/clustalw]: . .
- Massari ME, Grant P, Pray-Grant MG, Berger SL, Workman JL, Murre C: A conserved motif present in a class of helix-loop-helix

proteins activates transcription by direct recruitment of the SAGA complex. *Mol Cell* 1999, **4:**63-73.

- Aronheim A, Shiran R, Rosen A, Walker MD: The E2A gene product contains two separable and functionally distinct transcription activation domains. Proc Natl Acad Sci USA 1993, 90:8063-8067.
- Kho CJ, Huggins GS, Endege WO, Hsieh CM, Lee ME, Haber E: Degradation of E2A proteins through a ubiquitin-conjugating enzyme, UbcE2A. J Biol Chem 1997, 272:3845-3851.
- 19. Loveys DA, Streiff MB, Schaefer TS, Kato GJ: The mUBC9 murine ubiquitin conjugating enzyme interacts with the E2A transcription factor. *Gene* 1997, 201:169-177.
- 20. [http://genome.ucsc.edu/index.html]: . .
- Meyer KB, Mufti DAH: Post-transcriptional regulation of E2A proteins via lipopolysaccharide and CD40 signaling. Eur J Immunol 2000, 30:719-724.
- 22. Sigvardsson M, O'Riordan M, Grosschedl R: **EBF and E47 collabo**rate to induce expression of the endogenous immunoglobulin surrogate light chain genes. *Immunity* 1997, **7:**25-36.
- 23. Benezra R, Davis RL, Lockshon D, Turner DL, Weintraub H: The protein Id: a negative regulator of helix-loop-helix DNA binding proteins. Cell 1990, 61:49-59.
- Christy BA, Sanders LK, Lau LF, Copeland NG, Jenkins NA, Nathans D: An Id-related helix-loop-helix protein encoded by a growth factor-inducible gene. Proc Natl Acad Sci USA 1991, 88:1815-1819.
- Kee Y, Bronner-Fraser M: Id4 expression and its relationship to other Id genes during avian embryonic development. Mech Dev 2001, 109:341-345.
- Parvari R, Ziv E, Lantner F, Heller D, Schechter I: Somatic diversification of chicken immunoglobulin light chains by point mutations. Proc Natl Acad Sci U S A 1990, 87:3072-3076.
- 27. Sale JE, Calandrini DM, Takada M, Takeda S, Neuberger MS: Ablation of XRCC2/3 transforms immunoglobulin V gene conversion into somatic hypermutation. *Nature* 2001, **412**:921-926.
- Meyer KB, Skogberg M, Margenfeld C, Ireland J, Pettersson S: Repression of the immunoglobulin heavy chain 3' enhancer by helix-loop-helix protein Id3 via a functionally important E47/E12 biding site: implications for developmental control of enhancer function. Eur J Immunol 1995, 25:1770-1777.
- 29. Quong MW, Massari ME, Zwart R, C. Murre.: A new transcriptional activation motif restricted to a class of helix-loop-helix proteins is functionally conserved in both yeast and mammalian cells. *Mol Cell Biol* 1993, **13:**792-800.

