Modifying inter-cistronic sequence significantly enhances IRES dependent second gene expression in bicistronic vector: Construction of optimised cassette for gene therapy of familial hypercholesterolemia

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ABSTRACT

Internal ribosome entry site (IRES) sequences have become a valuable tool in the construction of gene transfer and therapeutic vectors for multi-cistronic gene expression from a single mRNA transcript. The optimal conditions for effective use of this sequence to construct a functional expression vector are not precisely defined but it is generally assumed that the internal ribosome entry site dependent expression of the second gene in such as cassette is less efficient than the cap-dependent expression of the first gene. Mainly tailoring inter-cistronic sequence significantly enhances IRES dependent second gene expression in bicistronic vector further in construction of optimised cassette for gene therapy of familial hypercholesterolemia. We tailored the size of the inter-cistronic spacer sequence at the 5′ region of the internal ribosome entry site sequence using sequential deletions and demonstrated that the expression of the 3′ gene can be significantly increased to similar levels as the cap-dependent expression of the first gene. The inter-cistronic spacer sequence at the 5′ region of the internal ribosome entry site sequence involving direct RNA-RNA contacts and RNA-protein interactions. These results provide a mechanistic basis for translation stimulation and RNA resembling for the synergistic stimulation of cap-dependent translation.

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1. Introduction

Simultaneous expression of multiple heterologous genes from a single gene transfer vector is an important requirement in a range of gene transfer and gene therapy protocols. There are generally four common strategies in which two or more genes can be co-expressed from a single vector. Firstly, two genes can be fused together in frame to produce a chimeric sequence ensuring simultaneous expression of both genes as a fusion protein [1–3]. However, this strategy may not work for all fusion proteins as some may be misfolded or mistargeted and, particularly important to gene therapy, the fusions might be immunogenic to the host. Secondly, separate promoters can be used to drive expression of different genes in the same vector. The major disadvantage of constructing such a dual promoter vector is possible transcriptional interference and/or dissociated gene expression, with a fraction of the transfected or transduced cells expressing only one gene such as the selectable marker and not the gene of interest and vice versa [4–6]. The third strategy relies on the utilisation of natural splicing signals of viruses by which multiple RNAs are produced from a single transcript [7]. However, this is not a frequently used strategy because of sub-optimal splicing of the viral vector. To overcome the above shortcomings, the fourth strategy involves the construction of a bicistronic

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**Fig. 1.** Structures of the bicistronic cassettes. The sequence and minimum energy structure of the IRES (Emcv) highlight motifs of interest for bicistronic protein expression. RNA sequences near the IRES AUG differed for commercial and derived plasmids. Uppercase denotes native Emcv bases. The protein coding ORFs and all plasmids encode A6 in the JK bifurcation loop except pCITE-1, pIREs, and pF/R-att. A) Schematic diagram of the constructed 5'-LDLR-IRES-eGFP-3' expression cassettes with sequential deletions at the 5' of IRES element. Sequence annotation of spacer sequence in between IRES (Emcv)-eGFP. B) Inter-cistronic spacer sequence with different construct vectors.
bicistronic cassettes, in which the two heterologous genes are separated by an element known as the internal ribosome entry site (IRES) sequence. Transcription of both genes in the bicistronic cassette is driven from a common upstream promoter, thus eliminating promoter interference. As a result, a single messenger unit including the bicistronic transcript of both genes separated by IRES is produced to allow uncoupling of the two Cistrons during translation. Translation initiation of the first Cistron 5′ from the IRES is typically mediated by cap-dependent translation initiation [8], however translation of the second Cistron 3′ from the IRES is mediated independently of capping by using the IRES element as an internal ribosome-binding site to initiate translation.

Since both the first and the second genes in a bicistronic cassette are under the control of the 5′ upstream promoter, detection of the protein encoded by the second Cistron is theoretically assured by the successful expression of the first Cistron. However, many researchers using IRES elements for construction of bicistronic gene transfer vectors have reported difficulties in achieving significant expression of the second Cistron. Studies conducted to determine the parameters that influence IRES-dependent translation initiation in bicistronic constructs have shown that the length and the secondary structure of non-coding inter-Cistronic sequences (ICS) that separate the 3′ end of the IRES element from the downstream Cistron have a strong effect on the efficiency of IRES function [9–11]. However, little is known about the ideal length of non-coding ICS that separates the 5′ end of the IRES element from the upstream Cistron and provides optimal expression both 5′ and 3′ of the genes.

In the present study, we constructed nine expression vectors containing bicistronic cassettes under control of the CMV promoter. These bicistronic cassettes contain sequentially deleted non-coding ICS sequences, between the stop codon of the human low-density lipoprotein cDNA (hLDLR cDNA) and the IRES-eGFP cassette sequences. We then examined the efficiency of these vectors with the aim to screen and select for an optimal 5′-hLDLR-IRES-eGFP-3′ cassette, which can be used for preclinical gene therapy studies of familial hypercholesterolemia.

IRES direct recruitment of the initiation complex without requiring a 5-cap [12]. Generally present in the 5-UTR, IRES-dependent initiation may occur by direct binding of the 4OS subunit to the mRNA, or may be mediated by a subset of the canonical initiation factors, and in some cases necessitates specific cellular proteins called IRES trans-acting factors (ITAFs) [13]. Furthermore, in silico analysis of RNA structure prediction, comparative sequence analysis and minimization of thermodynamic free energy equilibrium structure were applied with our IRES search system.

To better understand the structures and functions of IRES RNAs. We used comparative sequence analysis (CSA) to analysis and identity covering base pair. This approach was proven to be effective in the construction of reliable secondary structure models of ribosomal, signal recognition particle and transfer messenger RNAs [14]. Our studies yielded a revised secondary structure of IRES RNA that was supported by both covering base pairs and available biochemical data. The model was used to investigate IRES RNA in three dimensions in free form as well as when bound to IRES associated proteins and the 40s ribosomal subunit.

2. Results

2.1. Structure of the bicistronic cassettes

In view of the less dramatic effect of inter cistronic sequence (ICS) on IRES mediated translation initiation, to test the effect of ICS length on gene expression we constructed nine expression vectors containing bicistronic cassettes with variable length of ICS sequences (Fig. 1A and B). Because the order of the genes on the bicistronic cassette has a great influence on their expression levels [9] we placed the hLDLRcDNA sequence that we intend to use as a therapeutic gene for functional complementation of hLDLR deficiency, 5′ to the IRES element. Sequential deletions were created using a modified ExoIII/S1 nuclease protocol [15] and the DNA fragments that were generated containing the hLDLRcDNA with different length of spacers were subcloned into the pIRES2-eGFP (Clontech) mammalian expression vector. The pIRES2-eGFP expression vector was chosen because it contains the IRES of the encephalomyocarditis virus (EmcV) which has a high translation efficiency compared to other IRES sequences including those from hepatitis A and C viruses, poliovirus, human rhinovirus, and foot-and-mouth disease virus [16–19]. In addition to its high efficiency, EmcV IRES posses a broad tissue tropism [20] which makes it the most widely, used in gene therapy and gene transfer protocols [21–23]. This vector also permits high levels of plasmid DNA production and its multi-cloning sites are helpful for gene cloning. The eGFP incorporated in this vector is a red-shifted variant [24,25] of wild type eGFP, which has been optimised for brighter fluorescence and higher expression in mammalian cells.

Moreover, sequences flanking the eGFP have been converted to carry a Kozak consensus translation initiation site [26] that increases further translation efficiency in eukaryotic cells. The AUG initiator codon of the eGFP is located 12 bases downstream of the IRES element, i.e., such a short distance is nevertheless sufficient to significantly affect the eGFP expression, regardless of the conservation of the optimal eGFP AUG context (acc AUG gug). Indeed, flow cytometry analyses have previously shown a more than 10 fold difference in eGFP peak intensity.

The eGFP was used as a marker for clear and rapid detection for monitoring gene expression. As the fluorescence activity for eGFP requires IRES no substrates, cofactors or additional gene products the transfected cells can be sorted immediately by flow cytometry without the use of antibiotic for selection when such a construct is used for future in vivo studies.

2.2. Analyses of hLDLR and eGFP expression by western blotting and flow cytometry

Comparison of expression from the first and the second gene between cells transfected with bicistronic constructs containing variable length of spacer was performed by Western blotting and flow cytometry. Western analysis ensures that the expressed proteins are of the correct, mature sizes and the flow cytometry permits accurate assessment of gene transfer and expression at a single cell level. Immunoprophbing with an anti-hLDLR antibody (Fig. 2A) shows the mature and immature forms of the hLDLR protein from cell extracts transfected with each of the nine constructs. The hLDLR migrates as two bands of molecular masses of 160 kDa and 120 kDa of which the 160 kDa mature band was more prominent in all samples. No signals were detected from the negative controls (lane 10, 12, and 13). As expected, the level of hLDLR protein from cells transfected with the mono-Cistronic pLDLR-C3 vector (lane 11) was higher than the amount of hLDLR protein produced from all other bicistronic cassettes. To generate quantitative data for comparative analysis of these blots, each signal on the blots was measured by densitometry and normalised against β-tubuline (Fig. 2B and C) using National Institutes of Health (NIH) software by ImageJ, a Java program [27]. No significant differences in hLDLR expression were detected from samples transfected with constructs containing different spacer sizes. Immunoprophbing of the membrane with anti-eGFP antibody in Fig. 2B shows the production of eGFP protein of the expected molecular mass of approximately 30 kDa from cells transfected with clones containing ICS sequences ranging from 188bp to 18bp. Unlike hLDLR expression, the levels of eGFP expression increase with decreasing length of the spacer sequence, which is also confirmed by densitometry after normalisation with β-tubuline (Fig. 2D). The highest bicistronic level of eGFP expression is detected in extracts from cells transfected with a construct containing a 34bp spacer (lane 8). The expression decreased somewhat again when the spacer is shorten to 18bp (lane 9).
activity in the spacer sequences which might promotes eGFP expression analyses should exclude the presence of a putative intrinsic promoter lengths were performed using Northern blotting and qRT-PCR. These transfected with bicistronic constructs containing variable spacer sequences between the stop codon of hLDLRcDNA and the IRES element summary, the data provided above show that the length of the ICS sequence. The eGFP signals were undetectable from cells transfected with constructs containing spacers of less than 216bp. The levels of eGFP expression were, generally, increasing with decreasing length of the spacer sequence. The eGFP signals were undetectable in cells transfected with constructs IRES sizes: (1, 445bp, 2, 372bp, 3, 247bp, and 4, 216bp). D) Quantitative densitometry analysis of protein expressions of LDLR and eGFP.

However, no eGFP signals are detectable from cells transfected with constructs with spacer lengths of 445bp, 372bp, 247bp, and 216bp (lane 1–4). In contrast, the amount of eGFP protein from cells transfected with the mono-Cistronic peGFP-C3™ plasmids (lane 12) is higher than the amount of eGFP protein produced from all other bicistronic cassettes. It is noteworthy that mono-Cistronic cassette containing IRES-eGFP mother cassette (lane 10) generates low eGFP protein compared to peGFP-C3 mono-Cistronic vector (lane 12) and also to bicistronic vectors containing 18bp, 34bp 113bp, 141bp and 188bp spacers (lane 5–9). No other proteins were recognised from the cell extracts immunoprobed with these antibodies. Flow cytometry of the transfected cells (Fig. 3 A & 3B) show the mean fluorescence intensity and percentage of eGFP positive cells, which confirm the data obtained by Western blotting. Expression of eGFP is almost undetectable from cells transfected with constructs with ICS lengths of 445bp, 372bp and barely detectable from cells transfected with constructs with spacer lengths 247bp, and 216bp. The expression of eGFP peaks at a spacer length of 34bp and starts to decrease when the spacer is shortened to 18bp. In summary, the data provided above show that the length of the ICS sequences between the stop codon of hLDLRcDNA and the IRES element correlate directly with the levels of expression of the second Cistron independently. Northern analysis should also ensure that the expressed proteins are generated from the correct sizes of bicistronic mRNA transcripts. Northern blot analysis with the PCR generated hLDLRcDNA probe (Fig. 4A) revealed production of the correct ~ 4 kb 5’-LDLR-IRES-eGFP-3′ single mRNA transcript from cells transfected with constructs vector 6 to 9 containing 141bp, 113bp, 34bp, and 18bp ICS sequences. Indeed, this blot indicates that both hLDLR and eGFP genes from the respective bicistronic cassettes were transcribed as one transcriptional unit and independently translated into two proteins. In contrast, cells transfected with bicistronic cassettes 1 to 4 with ICS of 445bp, 372bp, 247bp, and 216bp, showed the production of the 2.6 kb hLDLR mRNA transcripts only. Unexpectedly, the amount of the 2.6 kb hLDLR mRNA transcripts produced from these bicistronic cassettes was as high as from the control mono-Cistronic hLDLR construct vector 12 compared to those with shorter ICS sequence. This indicates that the long spacer might function as a transcriptional termination signal leading to the generation of mono-Cistronic transcripts instead of the bicistronic one.

Although, the Cistron located at the 5′ of IRES is preferentially produced and at a high levels. Unexpectedly, cells transfected with bicistronic cassette with spacer of 188bp (construct 5) showed the production of both the 2.6 kb hLDLR mRNA and the 4 kb 5′-LDLR-IRES-eGFP-3′ mRNA transcripts. This suggest that at this spacer length, transcription termination might occur prematurely, although, inefficiently leading to the generation of the two different sizes of transcripts. We next ruled out the presence of a putative intrinsic promoter activity in the constructs or the spacer sequences which might promotes eGFP expression independently by blotting the membrane using the PCR generated eGFP probe (Fig. 4B). As expected, the correct ~ 4 kb 5′-LDLR-IRES-eGFP-3′ single mRNA transcript from cells transfected with constructs 5 to 9 containing 188bp, 141bp, 113bp, 34bp, and 18bp ICS was detected. No signals were detected from cells transfected with constructs 1 to 4 containing 216bp, 247bp, 372bp, and 445bp spacers. In addition, no eGFP mono-Cistronic mRNA signal was detected from any of the constructs, which excludes the presence of a putative
Fig. 3. Flow cytometry analysis of IRES-mediated eGFP expression from various bicistronic constructs. A) 10000 events were recorded and the percentage of positive cells (given as % values on histograms) and the geometric mean of eGFP fluorescence (given as numbers on histograms) were calculated using Cell Quest software. B) Bar diagram showing expression of eGFP in various constructs.
In order to test the functionality of the LDLR-IRES-eGFP-8 optimised cassette, we transfected the LDLR-deficient CHOIdlA7 cells and confirmed correct localisation of both the hLDLR and the eGFP transgenic proteins. We next investigated the ability of the transgenic hLDLR to bind and internalise fluorescently labelled-LDL (Dil-LDL™) in vitro. Living CHOIdlA7 cells were transfected with the pLDLR-IRES-eGFP-8 vector and assayed microscopically (Fig. 6). Only CHOIdlA7 cells which had been transfected with the pLDLR-IRES-eGFP-8 (panel A) and those transfected with pLDLR-C3 (panel B) expressing native LDLR were able to internalise Dil-labelled LDL particles. No internalisation of Dil-LDL™ is observed in non-eGFP expressing CHOIdlA7 cells shown on the same slides and CHOIdlA7 cells transfected with the peGFP-C3™ plasmid (panel C). Assuming that in vivo experiments are best suited to test the efficacy of the optimised 5'-hLDLR-IRES-eGFP-3' cassette, a pilot experiment was conducted by injecting 20 μg of plasmid DNA into adult MF1 mice fed a normal chow diet. Vector delivery was performed using hydrodynamic gene transfer method [1,2]. The peGFP-C3™ plasmid was injected as an eGFP control. Injected mice were sacrificed approximately 72 h after vector delivery and liver cells analysed microscopically confirmed eGFP-expressing cells (approximately 15–20% of liver mass).

2.5. Interaction between IRES RNA structure and 40S human ribosomal subunit

There was no structure available for IRES (Emcv). To the best of our knowledge, this is the first time we predict the IRES RNA structure model of all its domains and Emcv IRES RNA protein complex using homology modeling through MD simulation that's also showed an interactions with the 40S human ribosomal subunit during translation initiation (Fig. 7A–E). The topography of the initiation complex was assessed using the predicted protein domains ribosome and IRES: ribosome interactions as well as contact data information. It has been found that the interactions of elf4F and elf3 with type II IRES RNAs are the most useful [28,29]. These two aspects are essential for the recruitment of Emcv IRES RNAs to ribosomes [30]. Elf4F acts in conjunction with elf3 as a connection between the IRES RNA and the 40S human ribosomal subunit. The C-terminal portion holds the binding sites for both the IRES RNA as well as elf3 [31]. It binds to the 40S human ribosomal subunit in the absence of other initiation factors and has been shown to make multiple contacts with the Emcv IRES RNA [32]. Although the exact location of the elf3 binding sites on Emcv IRES RNA is not clear, Cryo-EM studies of elf4F- G: elf3 and elf3: 40S complexes place the Emcv-associated elf4F near the ribosomal E site [33]. Due to the lack of data for this complex near the E site of the ribosome, Emcv IRES: elf4F interactions were not modeled (Fig. 7E), consistent with the models of elf4F: elf3 and elf3: 40S complexes.

2.4. Analyses of biological activity and in vivo expression of optimised cassette

In order to test the functionality of the LDLR-IRES-eGFP-8 optimised cassette, we transfected the LDLR-deficient CHOIdlA7 cells and confirmed correct localisation of both the hLDLR and the eGFP transgenic proteins. We next investigated the ability of the transgenic hLDLR to bind and internalise fluorescently labelled-LDL (Dil-LDL™) in vitro. Living CHOIdlA7 cells were transfected with the pLDLR-IRES-eGFP-8 vector and assayed microscopically (Fig. 6). Only CHOIdlA7 cells which had been transfected with the pLDLR-IRES-eGFP-8 (panel A) and those transfected with pLDLR-C3 (panel B) expressing native LDLR were able to internalise Dil-labelled LDL particles. No internalisation of Dil-LDL™ is observed in non-eGFP expressing CHOIdlA7 cells shown on the same slides and CHOIdlA7 cells transfected with the peGFP-C3™ plasmid (panel C). Assuming that in vivo experiments are best suited to test the efficacy of the optimised 5'-hLDLR-IRES-eGFP-3' cassette, a pilot experiment was conducted by injecting 20 μg of plasmid DNA into adult MF1 mice fed a normal chow diet. Vector delivery was performed using hydrodynamic gene transfer method [1,2]. The peGFP-C3™ plasmid was injected as an eGFP control. Injected mice were sacrificed approximately 72 h after vector delivery and liver cells analysed microscopically confirmed eGFP-expressing cells (approximately 15–20% of liver mass).

The intramolecular interaction in domains of IRES (Emcv), showed
that the specific binding receptor of in tetra loop was unknown, although the tertiary contact was required for efficient IRES activity, the specific binding receptor of in tetra loop was unknown. We selected a final Model IRES (Emcv) and found two receptor candidates (G231/C241, and C232/G240) base pairs interacting with A180 and A181 residues in GUAA tetra loop. The trajectory for IRES (Emcv) shows that the two adenosines retained a distance <3 Å. In contrast, only the first adenosine A180 of other Models retain a distance <4 Å during the initial 10, and 20 ns, respectively. In Model IRES (Emcv), the average distance between C232/G240 pair and A180 is 2.0 ± 0.50 Å while C231/G241 pair and A181 was 2.0 ± 0.21 Å. These findings targeted receptors of A180 and A181 residues, respectively. The tertiary interaction of IRES (Emcv), we considered the Leontis/Westhof nomenclature and analysed the three edges Watson Crick, Hoogsteen and Sugar edge for potential hydrogen bonding interactions. The measured minimum distances between the Sugar edge of each C232 to G240 and C231 to G241 base pair with three edges of each A180 and A181 over the 100 ns time showed tightly formed hydrogen bonding interactions for the sugar edge/Watson–Crick between the C232/G240 pair and A180 and sugar edge/Hoogsteen edge tertiary interactions between G231/ C241 pair and A181. In addition, we observed tertiary contacts between U179 and A234 residues via trans Watson–Crick/Watson–Crick edge interactions at 22 ns? These long-range interactions occurred sequentially at 7, 20 and 22 ns, involving A180, A181 and U179, respectively. These co-operative long-range interactions helped in the stabilization of the IRES domain.

3. Nine constructs of spacer sequence and thermodynamic free energy

IRES calculate spacer sequence of 1–9 constructs minimum free energy of the thermodynamic ensemble mRNAs structures. The genetic code of the sequences introduced has to be in the correct form. It was necessary then to change all the timelines present in the sequences to the uracile residues. The optimal secondary structure symbolized with brackets and points, and it also shows the minimum free energy in kcal/mol of each IRES spacer sequence construct structure (Fig. 8). The more negative results are more stable thermodynamically equilibrium structure. The secondary structure predicted by RNA-Composer is due to the pattern predicted structures corresponded to the real IRES structure of our target. The pattern we had only used one nucleotide upstream the loop-bulge. For this reason, the secondary structure was not as stable as if more base pairings had been included in the stem. In all nine IRES constructs predict the molecular dynamic of tertiary structure shown in the clear thermodynamic energy scale level. The enthalpy and entropy are state functions, so changes in their values depend on the path between as initial and final state. We did for the free energy construct thermodynamic cycles for the changes in enthalpy and entropy for the good free energies and stability of the IRES spacer sequence construct.

4. Discussion

When an IRES is used for the construction of a multi-cistronic cassette in gene transfer and expression vectors, it has been frequently observed that the DNA sequence subcloned at the 3′ to the IRES element

![Image](https://via.placeholder.com/150.png?text=Fig. 6. Internalisation of Dil-labelled LDL by living transfected CHOIdA7 over-expressing the optimised 5′-hLDLR-IRES-eGFP-3′ cassette. Cells on panel A to C were transfected with pLDLR-IRES-eGFP-8, pLDLR-C3, and pEGFP-C3™ respectively. Only eGFP expressing cells on panel A or those expressing native LDLR (panel B) were able to internalise Dil-labelled LDL particles. No internalisation was detected in non-eGFP expressing cells on same slides or in those transfected with the pEGFP-C3™ plasmid (panel c). PH is phase contrast.)
is poorly expressed compared to the sequence subcloned at the 5’ of the IRES [34]. In most of the reported articles this phenomenon is attributed to the low efficiency of IRES mediated translation initiation compared to cap dependent translation. A possible explanation is a higher affinity of translation factors for the cap structure than for the IRES element. Therefore, transcription factors may become less available for the internal initiation of translation [35]. Alternatively, the IRES may require additional host trans-acting factors, which modulate its function in a cell or tissue type specific manner [36]. Thus, high levels of IRES mediated internal translation initiation can be
Fig. 7. Emcv (IRES) structural and functional annotation. A) Three-dimensional model structures of IRES (Emcv) RNA domains modeling structures consists in to two form of (domains Residues. 273–697); D, E, IA, IB, IC, GA, GB, H, F) is represent in the helical form. B) The complete domains of IRES 2D structure of the RNA with highlighting in grey Color for all domains. C) Other (domains Residues. 680–836; J, K, L, M). Domains are in different colors as in the secondary structure schematic depicted at the lower right. The 5’ and 3’ ends of domains along with conserved and variable subdomains are shown. The main helical stalks are shown in orange and yellow. The locations of helices and the tetra loop are shown in red. Bifurcation loop residues represent in red Color. The images were generated using QTMG. X-ray crystal structure of eGFP start coding with MET and 12bp downstream in (red) of the IRES element were representing different colors blue are AUG stop codon and initiate of start codon eGFP MET. D) Secondary structure of IRES (Emcv) RNA interaction for 40S human ribosome subunit. The interacting residues are highlighted in red Color. E) Hypothetical placement of protein-constrained IRES (Emcv) RNA on the Cryo-EM surface representing of the human 40S subunit. Cryo-EM reconstruction of the 40S subunit portion of the human ribosome with traveling of mRNA 5’ to 3’ (entry to exit channel) and also showing interaction with IRES (Emcv) entry in the APE site. Coloring of the IRES (Emcv) RNA ribbon colors were in green (J), blue (K), yellow (L) and orange (M). The first start codon (green) is indicated, and the naked human 40S subunit (EMDB ID 1092) was retrieved from the EM-Database. These images were generated using UCSF Chimera.

Fig. 8. A constructs of 1–9 spacer sequence and free energies. The different thermodynamic ensemble free energy of all constructs [1–9] of spacer sequence and the molecular dynamic structure were determined by simulation. A) The 2D secondary structure of EMCV-IRES constructs (C1–C9) as shown in RNA 2D view black. B) The secondary structure of EMCV-IRES in 1–9 as shown in red bar with denoted different thermodynamic free energy. C) The tertiary EMCV-IRES structures as shown in the helical state.
anticipated depending on the nature of the IRES sequence and its tissue tropism [16]. Another important factor that might have a strong influence on IRES dependent translation initiation is the length and possibly the nature of the ICS sequence flanking the IRES element at both its 5' and 3' ends [9].

In the present investigation, an IRES derived from the Emcv genome was used to mediate eGFP marker gene translation initiation. We have successfully constructed nine bicistronic cassettes containing sequentially deleted non-coding ICS sequence, between the stop codon of the hLDLRcDNA and the IRES/eGFP elements. We then analysed the expression levels of the respective transgenes systematically at the DNA, RNA and protein levels. By tailoring the size of the ICS sequence at the 5' of IRES, we showed for the first time that IRES dependent expression of the second gene could be significantly increased to the level of the cap-dependent expression of the first gene. Maximum expression efficiency of the eGFP was obtained when the ICS was 18bp to 141bp in length. However, when the ICS was 216bp or longer, the expression levels of eGFP were reduced significantly and become completely undetectable at a spacer length of 445bp. These results are in agreement with data published by Attal et al. (1999a) who found that IRES elements from Emcv as well as from poliovirus function optimally when about 100 nucleotides were added after the termination codon of the first Cistron (luciferase). These authors also showed that IRES elements become totally inefficient when added after nucleotide (300–500) spacer [9]. In their constructs the second Cistron was the chlor-amphenicol acetyl transferase gene (CAT). In a similar study using the IRES elements from poliovirus and from SV40, respectively in bicistronic cassettes containing the firefly luciferase gene as the first Cistron and the CAT gene as the second Cistron, Attal et al. (1999b) also showed that maximum efficiency in the expression of the second Cistron was achieved when the spacer fragment was composed of 30–90 nucleotides. This expression was profoundly reduced when the ICS fragment contained only 8bp and also when it was 300bp long. Finally, expression was undetectable when the spacer fragment was 500 nucleotides [37]. However, despite the remarkable strong influence of the length of ICS sequence on gene expression of the second Cistron in Attal studies, the levels of expression from the second gene were never similar or exceeding those of the first gene.

It has been frequently argued that the low level of second Cistron expression is due to the low efficiency of IRES functions in the cell in questions. We showed here that the low levels of expression from the second gene when ICS of 218bp or longer is used, is due to the generation of a single transcriptional unit containing the LDLR Cistron only, suggesting that long ICS sequence may serve as an insulator for transcription termination. To exclude the effect of tissue tropism, we have transfected our constructs into Hela and CHOIdlA7 cells and obtained similar results. We have also noticed that the amount of mRNA containing the hLDLR and eGFP Cistrons does not correlate tightly with the levels of translation of both Cistrons. We therefore cannot exclude that the sequentially deleted sequences might also contain an element that influences mRNA instability. However, we do not have detailed data to support this assumption, which is nevertheless worth further investigation.

To determine whether the phenomenon illustrated above can be generalized to other genes in this arrangement, expression from first and second gene was also compared using a bicistronic plasmid with hLDLRcDNA 5’ from a 445bp ICS identical to construct 1 and the Hygromycin gene in 3’ position. Hygromycin activity in HEK 293, Hela and CHOIdlA7 cells transfected by pLDLR-IRES-Hyg was undetectable. We have also constructed another bicistronic cassette with the human Olig2cDNA 5’ and the dsRed2 gene 3’ to a different 400bp ICS sequence. The fluorescent of dsRed2 protein was undetectable in HEK 293, Hela and human fetal mesenchymal stem cells transfected by this construct. Shorting the spacer by 200bp permits high expression of dsRed protein without affecting the expression level of Olig2 transcription factor. We are also aware that permanent phenotypic corrections of LDLR deficiency by gene therapy may require long-lasting expression of the transgenic LDLR in the relevant animal model to judge its efficiency, therefore, we have constructed and examined several viral plasmids based on the lentiviral backbone under the control of ubiquitous CMV promoter. However, the production of viral particles in HEK 293T producer cells following a standard three-plasmid co-transfection protocol failed to generate infectious vector. Owing to the utilisation of a previously characterised liver specific promoter (LSP) [38], Kankkonen and colleagues demonstrated for the first time successful construction and production of high titre (1 × 10⁹ IU) third-generation HIV-1 based lentiviral vectors encoding rabbit LDLR [42].

Comparative sequence analysis [39] is the standard procedure for predicting of RNA secondary structure with an all-atom model. Over 99% base pairs accuracy can be obtained in ribosomal RNA secondary structures, predicted by comparative sequence analysis similar to the one exist in high-resolution crystal structures [40]. However, comparative sequence analysis required IRES a large number of homologous sequences from various databases. In the absence of necessary homologous sequences, minimum free energy IRES RNA structure prediction can be applied to predict the structure of a single RNA sequence with an average of 90% accuracy through molecular dynamics (MD) simulation [41]. This accuracy is sufficient to be used as a starting point to build an alignment for comparative sequence analysis [42] for the target IRES RNA sequence. Identify a specific structure at the 3’ border of the IRES that prevents the correct positioning of the AUG-MET initiation codon in the close vicinity of the ribosomal P site. Taken together, these results demonstrate the presence of an RNA determinant in IRES that allows ribosomal 60S subunit binding, although it is not sufficient to correctly position the AUG in the ribosomal P site. This allows us to propose a new model for initiation complex assembly on IRES and to refine the role-attributed stabilizing and promoting the conformational changes.

We predicted a model of IRES (Emcv) and determined that IRES RNA secondary structures, which is reliable method to model, the biologically meaningful tertiary structure [43]. The base-pairs information was utilized for molecular structure prediction using RNA-Composer, which facilitates RNA structures as well as three-dimensional model with form of RNA helical information [44], based on reliable algorithms. The machine translation principle and operates on the RNA FRBASE database acting as the dictionary relating to RNA secondary structure and tertiary structure elements. Furthermore, Covariation-modeling analysis was not limited to defining secondary structure but may also be used to identify possible long-range interactions [45]. Enzymatic and chemical probing of mutant IRES (Emcv) RNAs suggested that the GNRA tetra loop capping helix might form a tertiary contact with 240-GCACG-244 in the helix [34]. Our attempts to identify canonical RNA-RNA tertiary interactions for constraining the model were successful. We used data symbolizing Polypyrimidine tract binding protein (PTB) interactions of Emcv IRES RNAs. PTB enhanced the translation initiation directed by IRES RNAs of Emcv and hepatitis A virus (HAV) [46]. It has been showed that the Emcv dependence on PTB is conditional [47]. IRES-RNA self-folding structural element due to the intramolecular RNA–RNA interacting region has thus been suggested to contribute significantly to the IRES structural organization and stability of domain, and to the critical function of IRES activity [30]. IRES-mediated translation initiation is closely linked to structural organization in domain, specifically the apical region formed by two four-way junctions enabling the RNA–RNA intramolecular interactions. Thus, we focused on the apical region of IRES domain to decipher the spatial arrangement of the RNA fold that is a prerequisite essential step to understand the initiation mechanism of translation. The contact between the 40S human ribosome subunit and IRES RNA was visualized by Cryo-EM. Cross-linking of rpS5 to rpS25 has been suggested in between these proteins [48]. Recent X-ray diffraction studies of the structure of eukaryotic ribosomes have also confirmed this proximity
and indicate that rpS5 was located at the solvent side of the head of the 40S subunit [49]. We therefore proposed that helix1 domain was similar to these ribosomal proteins. The translation factors of eIF1 and eIF1A promoted scanning by inducing an open conformation of the ribosomal subunit. The IRES was expected to be outside the ribosome by the time the second AUG codon enters the P site.

A recent study found that the competition between HCV domain III and eIF3 for binding with 40S subunit may result in the reduction of 43S complex formation and may thus favor translation of HCV mRNAs [50]. We did not find any variants with changes within the eIF3 binding sites in domain III. However, the relevance of identified mutations, especially at position 183, for eIF3 interaction and translation cannot be excluded. The thermodynamic equilibrium method predicts the IRES RNA structure with the lowest equilibrium free energy, as the sum of independent contributions of stacked and base pair instem and loops [51]. Although Lu et al., 2009 refers that the Maximization of expected accuracy provides a better approximation than free energy minimization [52], in the selected IRES structure, thermodynamic and weight scoring scheme provides the best estimation of RNA 2D structure based on a minimum free energy calculations.

However, The AUG codon initiate of MET site selection for the main preference for initiation and dependent in part on the context of the AUG codon. In this case, eIF2 does not contribute in the formation of a ternary complex but, slightly, may it bind directly to the ribosomal A-site and facilitate binding of the Met-tRNAi to the ribosomal P site during translation initiation. EIF2 activity is not subject to direct regulation and consequently may not contribute to the control of mRNA translation.

Future work will aim to subclone our optimised 5′-hLDLR-IRES-eGFP-3′ cassette into a vector with liver specific promoter (LSP) promoter. The use of LSP to drive the expression of our optimised bicistronic cassette may permit physiological expression of LDLR, thus, avoiding the pathological intracellular lipid accumulation [53,54] that may damage the transduced cells in the relevant animal model. In addition, it should permit easy evaluation of vector bio-distribution following gene delivery.

5. Materials and methods

5.1. Plasmid construction

The hLDLRcDNA with the 347bp non-coding spacer sequence was excised from pBC-hLDLR by digestion with HindIII and SacI and ligated into HindIII/SacI digested pLmuts28™ (NEB, UK) to produce SplitmutshLDLR. The hLDLRcDNA and the 445bp downstream spacer sequences were derived from the pRSVh-LDLR plasmid. Using a novel PCR-based method, we were able to create several clones of SplitmutshLDLR with sequentially deleted sequences downstream from the LDLR open reading frame (LDLR-ORF) with a unique HindIII and KpnI restriction sites. After restriction of these clones with HindIII and KpnI, the fragments containing the LDLR-ORF and the non-coding “space” sequence downstream were blunt ended with Kl enow and then ligated into BglII/ BamHI digested and blunt ended pRES-2-eGFP™ expression vector. All expression vectors containing bicistronic cassettes were successfully constructed and their integrity confirmed by DNA sequence analyses. Expression of the transgenic proteins from these vectors is driven by the promotor cytomegalovirus (CMV). The integrity of the hLDLR-insert in the isolated pLDR-IRES-eGFP clones was examined by restriction digestion and DNA sequence analysis. The pLDR-C3 mono-Cistronic hLDLR expressing plasmid, which serves as a positive control for hLDLR expression, was constructed by removing the IRES-eGFP cassette from the pLDR-IRES-eGFP-1 construct using SalI and NdeI enzymes. The plasmid DNA was then blunt ended by Kl enow and relegation with T4 DNA ligase (NEB, UK).

5.2. Gene transfer into cultured cells

Human embryonic kidney epithelial cells (HEK 293) and LDLR-deficient Chinese hamster ovary (CHOIdA7) cells were cultured with Dulbecco’s modified Eagle’s medium (Invitrogen or Sigma, UK), supplemented with 10% fetal calf serum (FCS). All cells were incubated at 37°C with 5% CO2 in a humidified incubator. Cells were seeded into 15 cm² dishes at 60–70% confluence with a density of 4–6x10⁶ cells and were transfected the following day with 10 μg of the expression vector DNA using FuGene 6 (Roche, UK) according to the manufacturer’s instructions. Forty-eight hours later, expression of fluorescent proteins was examined with a fluorescence microscope. All transfection experiments were repeated at least three times with similar results.

5.3. Flow cytometry analysis

Flow cytometry analysis was carried out for the detection of cellular expression of eGFP using a FACSscan flow cytometer (Beckton-Dickinson, UK) with the FL1 detector channel. The data were acquired and analysed with CellQuest software (BD). 48 h post-transfection transfected HEK 293T cells were rinsed with PBS, then analysed for GFP expression. 10000 events were recorded. Untransfected and Mock transfected cells were used as controls. Results were presented as a percent of positive cells. Mean fluorescence intensity was used as an indicator of relative expression of eGFP on given cell population.

5.4. Preparation of total RNA and northern blot analysis

Total RNA was extracted with TRIzol based on the manufacturer’s instructions (Invitrogen, UK) and treated with DNase I (NEB, UK). Thirty micrograms total RNA from each sample were mixed with 2.5 parts of RNA loading buffer in a total volume of 35 μl, heated for 3 min at 100°C and chilled on ice. Samples were then separated on a 1% denaturing agarose-formaldehyde RNA gel prepared by microwave melting of 3g agarose (Invitrogen, UK) in 200mls DEPC-treated H₂O. Boiled agarose was allowed to cool to 60°C, then 56mls of 5× MOPS buffer and 50mls of 37% formaldehyde were added). After 4h of electrophoresis at 100 V, the gel was soaked in 0.05 M NaOH for 20–30min with gentle agitation, rinsed with DEPC-treated H₂O and soaked in 20 × SSC for 45min. Single-strand RNA was then transferred from the agarose-formaldehyde gel to a solid nylon membrane, Hybond-N+ by the capillary method. Gene images random prime labelling™ kit (Amersham, LIFE SCIENCE, UK) was used for labelling of 50 ng-purified PCR generated DNA probe following the manufacturer’s instructions. The blotted membrane was pre-soaked in 5 × SSC buffer before hybridisation and then placed in a hybridisation bag containing 50mls of pre-hybridisation solution for 60 min at 60 °C with gentle agitation. The labelled probe was then added into a fresh hybridisation buffer and hybridisation was carried out at 60°C for 18 h. Following hybridisation and washing, the gene images CDP-star detection™ kit (Amersham, LIFE SCIENCE, UK) was used for detection of hybrids according to the manufacturer’s instructions.

5.5. Reverse transcription (RT) and quantitative polymerase chain reaction (Q-PCR)

cDNA synthesis was accomplished with 1 μg of total RNA in the presence and absence of reverse transcriptase (RT). The final reaction mixture contained 1 μg RNA, 10 units RNAsin RNase inhibitor, 10 mM dNTP mix, 0.5 μg random primers (Promega, UK), 5 × reverse-transcription buffer, with (RT+) or without (RT-), 100 units M-MLV reverse transcriptase (Promega, UK), in a final volume of 20 μl. Quantitative expression analysis of the housekeeping gene encoding the ribosomal L19 protein gene (GenBank accession no. NM_000981) and the genes of interest Ldr (GenBank accession no. AB209409), eGFP and IRES sequence was determined by Q-PCR using an ABI PRISM™ 7700 Sequence...
Detector following the ABI protocol (Applied Biosystems, UK). The Q-PCR assays were run in triplicate on 96-well plates. Primers (5′-3′ sequences used for quantitative analysis are: L19-F 5′-gggaagggttcagcaacct-3′; L19-R 5′-tttgccgaggctgc-3′; LDLR-F3 5′-tgagttcatttgccaca-3′; LDLR-R3 5′-catcctccagactgaccatctg-3′; eGFP-F1 5′-gctgcgtgctggtctat-3′; IRES-R1 5′-ttccgggccctcacatt-3′ were designed according to real-time PCR requirements using the Primer Express 1.7 software. All primer sets were free of primer-dimer products. SYBR® Green was used in the Q-PCR assays. A standard curve using a control placentale cDNA pool was obtained in each assay. Thermal cycle parameters included an initial incubation at 50°C for 2 min for 1 cycle; Taq activation at 95°C for 10 min for 1 cycle; repetitive denaturation at 95°C for 15 s and annealing at 60°C for 1 min, for 40 cycles. Following completion of the quantitative amplification of the samples of interest, values were obtained and analysed using the Sequence Detector 1.7 software (Applied Biosystems, UK). All triplicate cycle threshold (Ct) values had to be within 1 Ct of each other and the correlation coefficient ($r^2$) above 0.98. The quantitative values for each triplicate were averaged and the relative expression of the genes of interest was determined by a ratio of their expression to that of the housekeeping gene for the same sample.

5.6. Western blot analysis of protein expression

Forty-eight hours after transfection, cells were harvested by scrubbing and re-suspended in 30µl/10⁶ cells ice-cold lysis buffer (10 mM Tris pH 7.5, 1 mM EDTA, 1% NONIDET P-40 (Sigma, UK)), protease inhibitor cocktail at indicated dilutions (Roche, UK). Cell suspensions were then centrifuged at 13,000 × g for 20 min at 4°C. Supernatant were re-centrifuged again at 40,000 × g for 45 min at 4°C. The protein content of cell lysate supernatants was assayed by the bicinchoninic acid (BCA) method using a commercial assay kit from Pierce with BSA as a standard. Five micrograms of protein of each lysate were combined with 30 µg of bovine serum albumin, boiled for 5 min and were then separated on a 13% or 7% SDS-PAGE gel for GFP and LDLR protein detection, respectively. Gels were then blotted onto PVDF membranes with a Hoefer apparatus (200 mA). After 1 h incubation at room temperature in blocking solution (5% dried skimmed milk in PBS-Tween-20 0.1%), membranes were incubated overnight at 4°C with specific primary antibodies, diluted as specified below with blocking solution. After three 5-min washes in PBS-Tween-20, blots were incubated at room temperature for 1 h with peroxides-conjugated goat anti-rabbit antibody (Abcam) or rabbit anti-mouse polyclonal antibody (Dako, UK) diluted 1:1000 in blocking solution. Following the final wash, detection on autoradiography hyper-films was performed after inducing a chemiluminescence reaction with the Amersham Super signal detection kit. Densitometric analysis was performed using ImageJ. Primary antibodies used in this study were anti-LDLR (polyclonal, 1:2000; Research Diagnostic, USA) and anti-GFP (polyclonal, 1:2000; abcam, UK) and anti-beta-tubulin (1:2000; Sigma, UK).

5.7. Emcv-IRES molecular modeling

The models were constructed using the sequence of the hLDLR-IRES (Emcv) RNA from (GenBank accession number X74312 positions 273–836). Its was consisted of different domains (starts from 5′ polyC residue 273, D, E, F, G2, G3, H, I3, J, K, L M (AUG-end of the Stop codon). The models of Emcv-IRES were determined by MD simulation methods. Briefly, the secondary structure information was used as input for RNA-Composer [53] installed on an IBM Blue Gene/L supercomputer. Based on the Structural Classification of RNA (SCOR) database [55] initial models were modified. The coordinates of appropriate structures were taken from the Protein Data Bank (PDB) [56] and were further analysed using CCP4/QTMG [57,58]. Experimental data obtained from the literature and the PDB were used to validate the modeled structure. Finally, bond angles and lengths were corrected to produce biologically feasible conformations.

5.8. Molecular dynamics simulations for hLDLR-IRES (Emcv)-eGFP region

Each system was solvated with the explicit TIP3P water model within a water box of dimension 10 Å on each surface. Simulations were performed while using the Amber force field [58–60] with Na ions to neutralize the system charge. We minimize the system in two steps, first over the water and ion molecules holding domains fixed and, second, with all constraints removed. The minimization was performed using the Powell conjugate gradient algorithm. The initial equilibrations were achieved over 60 ps at constant temperature (300K) and pressure (1atm). Pressure was maintained at 1atm using the Langevin piston method, with a piston period of 100fs, damping constant of 50fs and piston temperature of 300K. Temperature coupling was enforced by velocity ressagreement every 2ps. Both minimization and equilibration are performed using the NAMD program [61]. For the production run, we simulated a conventional MD trajectory for 100ns with the force field using the NAMD package. The system was simulated at constant temperature (300K) and volume using weakly coupled Langevin dynamics of non-hydrogen atoms, with a damping coefficient of $c = 10$ ps$^{-1}$ with a 2fs time step maintaining bonds to all hydrogen atoms rigid. Non-bonded interactions were truncated at 12 and 14 Å for van der Waals and electrostatic forces, respectively. Periodic boundary conditions were applied, and the particle mesh Ewald method was used to calculate electrostatic interactions. All simulations using the NAMD package were run on IBM Blue Gene/L supercomputer at the computational center for bioinformatics and genomics lab, Makkah.

6. Conclusion

We have presented a comparative analysis of protein expression patterns between different bicistronic cassettes containing variable length of ICS sequence. By using this approach to generate intervening ICS sequences of tailored size, we have overcome the previously repeatedly reported problem of insignificant second Cistron expression and have produced a functionally optimised 5′-hLDLR-IRES-eGFP-3′ expression cassette containing a 34bp spacer.

Author contributions

Conceived and designed the experiments: FA, ZA, MT, HM, MC, and SA. Analysed the data: AB, BB, SW, CC, MT, and WK. Wrote the paper: FA, and ZA.

Author disclosure statement

Authors have no conflict of interest.

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