Rapid proteasomal elimination of 3-hydroxy-3-methylglutaryl-CoA reductase by interferon-\(\gamma\) in primary macrophages requires endogenous 25-hydroxycholesterol synthesis

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**Abstract**

Interferons (IFNs) play a central role in immunity and emerging evidence suggests that IFN-signalling coordinates sterol biosynthesis in macrophages, via Sterol Regulatory Element-Binding Protein (SREBP) dependent and independent pathways. However, the precise mechanisms and kinetic steps by which IFN controls sterol biosynthesis are as yet not fully understood. Here, we elucidate the molecular circuitry governing how IFN controls the first regulated step in the mevalonate-sterol pathway, 3-hydroxy-3-methylglutaryl-CoA reductase (HMGCR), through the synthesis of 25-Hydroxycholesterol (25-HC) from cholesterol by the IFN-inducible Cholesterol-25-Hydroxylase (CH25H). We show for the first 30-min of IFN stimulation of macrophages the rate of \(\textit{de novo}\) synthesis of the \(Ch25h\) transcript is markedly increased but by 120-min becomes transcriptionally curtailed, coincident with induction of the Activating Transcription Factor 3 (ATF3) repressor. We demonstrate ATF3 induction by Toll-like receptors is strictly dependent on IFN-signalling. While the SREBP-pathway dependent rates of \(\textit{de novo}\) transcription of \(Hmgcr\) are relatively unchanged in the first 90-min of IFN treatment, we find HMGCR enzyme levels undergo a rapid proteasomal-mediated degradation, defining a previously unappreciated SREBP-independent mechanism for IFN-action. These events precede a sustained marked reduction in \(Hmgcr\) RNA levels involving SREBP-dependent mechanisms. We demonstrate that HMGCR proteasomal-degradation by IFN strictly requires the synthesis of endogenous 25-HC and functionally couples HMGCR to CH25H to coordinately suppress sterol biosynthesis. In conclusion, we quantitatively delineate proteomic and transcriptional levels of IFN-mediated control of HMGCR, providing a foundational framework for mathematically modelling the therapeutic outcome of immune-metabolic pathways.

**1. Introduction**

3-Hydroxy-3-methylglutaryl-CoA reductase (HMGCR) operates as an important regulated step enzyme in the mevalonate-sterol biosynthesis pathway, which is responsible for the production of cholesterol and isoprenoid products \([1,2]\). HMGCR is also an important therapeutic target for a class of inhibitory drugs, known as statins, which are prescribed to reduce cholesterol levels in the serum. Statin treatment reduces cholesterol but leads to a compensatory increase of HMGCR via the SREBP pathway (schematically shown in Fig. 6) and which also increases the level of low-density lipoprotein (LDL) receptors, leading to reduction in plasma...
cholesterol levels [1]. Given the importance of HMGCR in the biosynthesis of sterols, HMGCR is subject to stringent and multiple regulatory feedback mechanisms.

Regulation of HMGCR at the transcriptional level is exquisitely sensitive to feedback control mediated by the Golgi-nuclear shuttling of the master transcriptional regulator of the sterol pathway, SREBP2 [3–5]. Under conditions of high intra-cellular sterol levels, excess oxysterols and cholesterol bind to INSIG and SCAP proteins, respectively. This blocks transport of mature SREBP2 from the ER to the golgi, where cleavage of SREBP2 and release of the active transcription factor would normally take place [6–9]. Under sterol-limiting conditions, sterol binding to INSIG and SREBP cleavage activating protein (SCAP) does not occur and SCAP becomes free to escort SREBP2 from the ER to the golgi, where SREBP2 undergoes proteolytic cleavage [10–14] to generate an N-terminal domain of SREBP2 which is released into the nucleus to activate multiple sterol biosynthesis related genes, including Hmgcr [15].

Regulation of HMGCR at the post-transcriptional and post-translational level operates independently of the SREBP regulatory pathway and represents an important mechanism for regulating HMGCR in response to changing sterol levels. Post-transcriptional regulation is exerted by oxysterol-dependent degradation, via an INSIG dependent ER-associated protein degradation (ERAD) system, involving an ubiquitin-proteasome degradation pathway [16,17]. Under sterol depleted conditions, the half-life of HMGCR is approximately 12 h. However, in the presence of excess sterols, degradation of HMGCR by the ERAD mechanism results in a half-life of about 40 min [18]. This highly sensitive and responsive proteomic mechanism contrasts with the relatively slow transcriptional regulation mediated by the SREBP pathway which takes several hours to exert an effect on HMGCR protein levels [19].

It is well established that the oxysterol, 25-HC, which is synthesized from cholesterol by the enzyme CH25H, has a potent ability to block cholesterol biosynthesis through INSIG induced ubiquitination and proteasomal degradation of HMGCR [4,20,21] and also by blocking ER translocation of SREBP2 [22]. However, 25-HC has been shown not to play a major role in cholesterol homeostasis but instead has emerged to have key immune modulatory functions [1,23]. We have been the first to report that HMGCR levels and sterol biosynthesis are suppressed by interferon (IFN) and in infected macrophages the reduction of HMGCR is strictly dependent on IFN-induced signalling [24]. Moreover, in this connection 25-HC has also been shown to have a range of effects on immune related host defence systems, including suppressing the differentiation of human monocytes [25], decreasing immunoglobulin A production [26], promoting immune cell guidance [23], moderating inflammasome activity [27] and amplifying inflammatory signalling via AP-1 [28,29]. Park and Scott reported that the production of CH25H is stimulated by TLR activation in macrophages and DCs [29]. In later studies, we and other laboratories demonstrated that the IFN-mediated production of 25-HC has broad anti-viral and immune modulatory activities [27,28,30,31]. It has also been reported by many groups including our own studies that RNA-interference (RNAi) mediated knockdown of HMGCR protein levels and pharmaceutical inhibition of HMGCR activity can suppress viral infectivity and growth [24,32–34]. In addition, it is worth noting that HMGCR inhibitors (statins) also have immune modulatory effects [35]. Collectively, these findings point to an emerging role for HMGCR as a control point in the regulation of immunomodulatory and host defence mechanisms. However, the precise role of 25-HC in the immune system has not been fully defined. Moreover, roles for SREBP-independent mechanisms for anti-viral and anti-inflammatory effects have yet to be defined.

In this study, we examine further the regulation of HMGCR following early innate immune stimulation by IFN–γ. We report for the first time systematic quantitative analyses of the IFN regulatory circuit in macrophages for CH25H, its repressor ATF3, and how they couple to HMGCR, revealing a previously unappreciated role for the IFN–mediated degradation of HMGCR enzyme levels that is shown to be critically dependent on the de novo synthesis of 25-HC. Overall, this study contributes to the dissection of the mechanisms underlying the regulation of HMGCR as an emerging effector target during innate-immune-metabolic signalling and provides a foundation for future modelling of pathway feedback control.

2. Experimental

2.1. Reagents and media

25-HC and mevastatin were purchased from Sigma (Sigma-Aldrich, UK). MG132 (Z-Leu-Leu-Leu-al) (CAS number: 133407-82-6) was purchased from Cambridge Bioscience. Murine recombinant IFN-γ was purchased from Perbio Science or Life Technology (PAC4033). Medium A: DMEM/F12 with GlutaMAX (Gibco, Life technologies) supplemented with 10% (v/v) fetal bovine serum (FBS) (Gibco, Life technologies), 10% (v/v) 1929 containing colony-stimulating factor 1 (CSf1) and Penicillin/streptomycin (PS) (Gibco, Life technologies); Medium B: DMEM/F12 with GlutaMAX supplemented with 3% (v/v) lipo-protein depleted serum (LPDS) (Sigma), 10% (v/v) 1929 containing CSf1 and PS. The following antibodies were used: monoclonal mouse anti-HMGCR (C-1, Santa Cruz Biotechnology), rabbit anti-β-actin (4967, Cell Signaling) and rabbit anti-β-tubulin (ab6046, Abcam).

2.2. Primary bone marrow derived macrophage (BMDM) culture generation

Wild-type BMDMs were derived from the femur and tibia isolated from C57BL/6 mice as previously described [24,30]. Ch25H−/− BMDMs were derived from the femur and tibia isolated from B6.129S6-Ch25Hm−/−/J mice (The Jackson Laboratory, UK) and grown in Medium A. Ijb−/− mice were from the Institute of Animal Breeding and Genetics (Veterinary University of Vienna). All procedures were carried out under project and personal licences approved by the Secretary of State for the Home Office, under the United Kingdoms 1986 Animals (Scientific Procedures) Act and the Local Ethical Review Committee at Edinburgh University. All cultures are routinely tested to ensure they are free of mycoplasma and endotoxin contamination.

2.3. Experimental treatment conditions

BMDMs were grown in 24-well plates with Medium A. For the dose titration of mevastatin, cells were treated overnight with Medium B containing various concentrations of mevastatin. For the dose titration of 25-HC, cells were treated as the figure legend described. The cells were then harvested for western blot analysis. For time course experiments, Prior to the experiment, IFN-γ stock was diluted as described [24]. Culture medium was aspirated from 24–well plates and 1 mL of Medium C containing IFN–γ (5 ng/mL) was added to wells at 1.5-h, 4-h and 9-h time points, respectively. Meanwhile, a parallel experiment, where the cells were treated with 25-HC (2.5 μM), was performed at the same time. The cells were harvested for western blot analysis. For the measurement of HMGCR protein levels with proteasome inhibitor MG132, BMDMs were pre-treated with Medium B containing MG132 (20 μM) for an hour, and then treated with Medium C plus MG132 (20 μM) containing IFN–γ (5 ng/mL) or 25-HC (2.5 μM),
respectively. The cells were incubated at 37 °C for another 6 h and HMGCR levels were determined by western blot analysis.

2.4. qRT-PCR analysis

QIAGEN Rneasy Plus Mini Kit (QIAGEN, Germany) was used to purify total RNA. The qScript One-Step Fast qRT-PCR kit, Low ROX (Quanta BioSciences, USA), Taqman probe/primer Hmgcr, Atf3, Ch25h, Ifi1 and Gapdh (Life technologies, UK) were used for the qRT-PCR measurement. qRT-PCR was undertaken according to manufactures instructions.

2.5. Western blot analysis

After treatments, cells were lysed directly in wells by adding 100 µL of RIPA lysis buffer (9806, Cell Signaling) supplemented with protease cocktail (Complete Protease Inhibitor Cocktail Tablets, Roche) and 1 mM Phenylmethylene sulfonyl fluoride (Sigma). Plates were incubated on ice for 20 min. Protein concentrations of whole-cell lysate fractions were then determined using the BCA Protein Assay Reagent kit (Thermo Scientific) according to manufactures instructions. Prior to SDS–PAGE, whole-cell lysates were mixed with 2X Laemmli sample buffer (Sigma) and incubated at 50 °C for 10 min. Thereafter, an equal amount of protein was subjected to 8% SDS-PAGE, after which proteins were transferred to Hybond ECL Nitrocellulose membranes (Amersham). The membranes were blocked with 5% skimmed milk (Sigma), probed with specific primary antibodies overnight at 4 °C, washed with PBST and bands were visualised by Hyperfilm ECL (Amersham) or Odyssey Fc Dual-Mode Imaging System (Li-COR Biosciences) with Clarity™ Western ECL Substrate (BIO-RAD). Image Studio Lite (Li-COR Biosciences) was used to analyse the bands.

2.6. BMDM de novo RNA labelling, isolation and microarray analysis

2.6.1. Time course analysis of RNA synthesis and half-life in Control-treated or IFN-γ-treated BMDMs

To enable a simultaneous analysis of RNA synthesis, abundance and decay, BMDMs were control treated or treated with IFN-γ at a concentration of 5 ng/mL. These treatments were then followed by 16 successive cycles of 4-thiouridine addition to the BMDM cultures and transcriptional termination at 30 min intervals until the end of an 8-h time course. 4-Thiouridine (Sigma) labelling in these experiments was undertaken as described by Dölken et al. [36].

Total RNA was then isolated using an RNeasy Midi kit (Qiagen) according to manufactures instructions, quantitated using a Nanodrop (Thermo Scientific) and integrity was confirmed using an Agilent Bioanalyser (Agilent UK). Newly transcribed RNA (ntRNA) was then isolated and again quantitated using a Nanodrop.

2.6.2. RNA labelling for time-course microarray analysis

Processing of 100 ng of total RNA samples (8 × 1 h for control treated and 8 × 1 h for IFN-γ treated = 16 in total) and ntRNA samples (16 × 30 min for control treated and 16 × 30 min for IFN-γ treated = 32 in total) for hybridisation to Affymetrix Mouse Gene 1.0 ST arrays was undertaken according to manufactures instructions (Affymetrix). Hybridisation, washing, staining and scanning of the arrays were also undertaken following standard Affymetrix protocols.

2.6.3. Data analysis for microarray time-course

After scanning and data capture, open-source R based software “Biocductor” was used to implement all quality control and statistical analyses.

The RNA method was then used for normalisation, background correction, and probe set summation [37]. To reduce the dataset for further analysis and enrich for reliably detected genes, values for median and 2 × the Median Absolute Deviation (2 × MAD) for negative control probe sets were calculated for each array. Intensity values for each probe on each total RNA array were then compared with their array specific 2 × MAD value and were retained if the intensity was equal to or greater than this value. After filtering and removal of all designated Affymetrix control probesets, 12,472 probes were retained for statistical analyses.

Since conventional clustering algorithms do not typically assign significance to differential expression of two conditions over time (on an individual gene basis) the MaSigPro algorithm was used [38]. Using this method, 2086 (ntRNA) or 4482 (total RNA) probes with notable differential profiles of expression across the 8 h of IFN-γ stimulation were identified.

2.6.4. Microarray analysis of total, pre-existing and newly transcribed RNA in resting BMDMs

Using the protocol of Dölken et al. [36] for de novo RNA labelling and isolation, we used Affymetrix Gene STv1.0 microarrays to analyse total RNA abundance in resting BMDMs at t = 0 h and total, newly transcribed (labelled) and pre-existing (unlabelled) RNA at t = 1 h. This was completed in triplicate. To enrich for reliably detected genes, values for median and 2 × the Median Absolute Deviation (2 × MAD) for negative control probe-sets were calculated for each array. Intensity values for each probe on each total RNA array were then compared with their array specific 2 × MAD value and were retained if the intensity was equal to or greater than this value. After filtering and removal of all designated Affymetrix control probe-sets, 5826 probes were classified as detected on all arrays. Mean values for the total RNA, newly transcribed RNA and pre-existing RNA abundance measurements were then calculated from triplicate measurements and this data was used for the calculation of a median half-life in the resting BMDMs as described below.

2.6.5. Calculation of median transcript half-life in resting BMDMs

If the mean total RNA transcript abundance at the start of a 1 h time interval is T0, the mean total RNA transcript abundance at the end of the interval Tt, the mean newly-transcribed RNA transcript abundance Nt and the mean pre-existing RNA transcript abundance Pt, we should observe that Nt/Tt + Pt/Tt = 1. In practice, however, we observed some systematic disagreement with this equality. To maximise agreement, therefore, we introduced two normalisation constants, α and β, and rewrote the above equation as αNt/Tt + βPt/Tt = 1, as proposed in [39]. We then optimised α and β by linear regression and used the constants obtained in the calculation of half-lives for the time course. To estimate a half-life value for every transcript detected by the microarrays in this experiment, we calculated the values in two ways: (i) using newly transcriptional RNA transcript measurements (left equation) and (ii) using the pre-existing RNA transcript measurements (right equation). A mean of the 2 values obtained from these equations was then calculated.

\[
t_{1/2} = \frac{-55\ln 2}{\ln \left(\frac{\alpha}{\beta} \frac{\alpha + \beta}{\alpha}\right)}, \quad t_{1/2} = \frac{-55\ln 2}{\ln \left(\frac{\beta}{\alpha}\right)}
\]

In these calculations, we assumed a delay of 5 min before the initiation of 4-thiouridine incorporation into newly synthesised RNA. The resulting labelling period is, therefore, 55 min. Using
the above protocol, a median half-life value for the 5826 RNAs detected in these experiments was estimated. This was subsequently used to assist in the normalisation of half-life data from our time-course analysis of control treated or IFN-γ treated BMDMs (described in next section).

2.6.6. Estimation of transcript half-life in control or IFN-γ treated BMDMs

From our time course experiments we obtained data from 1 h intervals in which total and newly transcribed RNA had been analysed (150–210, 270–330, 330–390 and 390–450 min). This data was used to estimate half-life values for Hmgcr transcript in the control treated or IFN-γ treated BMDM data. Since we had chosen not to hybridise pre-existing RNA samples to microarrays in our 8-h time course experiment, we used the formulae above to optimise the value of τ such that the median half-life across all transcripts in the control and IFN-γ treated courses was the same as the value calculated for the resting BMDMs. The output of these calculations provided multiple, discrete half-life estimates for each transcript across the 8-h control or IFN-γ time course.

2.7. Software and statistical analysis

Microarray data analysis was performed by R [40]. The EC50 and IC50 values were calculated by Matlab [41] and Graphpad (GraphPad Software, Inc., La Jolla, CA, USA). Unpaired Student’s t tests were performed by Graphpad. The half-life of Hmgcr mRNA was calculated using Matlab. Adobe illustrator and Graphpad were used to assemble figures.

3. Results

3.1. Molecular circuitry for the rapid induction and inhibition kinetics of de novo RNA synthesis of Ch25h and Atf3 by IFN in primary macrophages.

In resting cells, latent transcription factors (named STATs) associated with the IFN-receptor reside in the cytoplasm in an inactive state and upon IFN binding its receptor, leads within seconds, to phosphorylated active forms that translocate, within minutes, to the nucleus, binding target promoters to transcriptionally active genes. Our previous studies have shown that IFN activation of BMDMs results in the increased biosynthesis and secretion of 25-HC as a result of a rapid STAT1 recruitment to the promoter of the Ch25h gene within the first hour of treatment [30]. The Ch25h gene encodes an enzyme which is responsible for catalysing the production of 25-HC from cholesterol. The timing and duration of 25-HC synthesis have important implications for its immune functions which are, in part, mediated through the metabolic and immune modulatory re-programming of BMDMs upon infection [42,43]. At the transcriptional level, ATF3 is a well known transcriptional repressor for TLIR driven pro-inflammatory cytokines [43,44] including the induction of Ch25h [45]. ATF3 has also been reported to be induced by TLIR-IFN signalling [46–48] and more recently has been found to repress the IFN-β promoter, establishing a negative feedback loop [49]. A schematic summary is shown in Fig. 6. While previous studies have clearly indicated a rapid up-regulation of Ch25h expression in response to IFN, the precise rate kinetics of IFN-γ elicited Ch25h mRNA and Atf3 de novo synthesis have not been determined. Furthermore, evidence for a direct functional requirement of IFN signalling in TLIR induction of ATF3 in BMDMs has not been shown. Hence, our initial experiments sought to quantitatively analyse de novo synthesis of Ch25h mRNA and its negative regulator Atf3 in IFN-γ stimulated BMDMs and in BMDMs genetically ablated for the TLIR-induction of IFN-β.

For these experiments, we performed an 8-h high-resolution, systematic microarray analysis of changes in RNA synthesis, abundance and decay in BMDM treated with 5 ng/ml IFN-γ. Fig. 1a shows how the de novo synthesis of Ch25h and Atf3 RNA altered during this temporal analysis. Ch25h RNA synthesis rate was up-regulated more than 4 fold in treated cells (relative to the control) during the first 30 min after the addition of the cytokine. Synthesis of Ch25h RNA then peaked 60–90 min after treatment and was subsequently down regulated, returning to basal transcriptional levels by 210–240 min (Fig. 1a). The kinetics of the observed transcriptional activation and repression provide an explanation for the observed plateau by 8 h for the amount of intracellular and secreted 25-HC as previously reported [30]. Notably, de novo synthesis of Atf3 RNA, the product of which is a repressor of Ch25h transcription, increased 60 min after the addition of IFN-γ (Fig. 1a). Atf3 RNA synthesis then peaked at 120 min, a time coincident with a marked reduction in Ch25h RNA synthesis. These experiments show a rapidly initiated, but apparently tightly controlled, transient up-regulation of Ch25h RNA synthesis during the first 4 h of IFN-γ treatment in BMDMs.

Next, we examined whether the induction of Atf3 by TLIR-stimulation depends on the presence of an intact IFN-β response in BMDMs. For these experiments, wild-type or Ifnb−/− BMDMs were treated with the TLIR-3 agonist, Poly (I:C) (10 μg/ml), and the abundance of Ifi1 (as a marker for STAT1 activation), Ch25h, and Atf3 RNA was analysed at 2, 4, 6, 8, 10 and 24 h after treatment. Fig. 1b shows how overall RNA abundance of the interferon-regulated gene Ifi1 varied in our time-course. In wild-type BMDMs, poly(I:C) treatment resulted in a rapid increase in Ifi1 RNA abundance during the first 2 h of the time-course. Notably, between 2 and 8 h the abundance of this transcript remained elevated and subsequently decreased from 10 h until 24 h. In Ifnb−/− BMDMs, in the absence of IFN-β production, the overall effect of the Poly(I:C) stimulus was negated. Alterations in Ch25h transcript expression elicited by Poly(I:C) followed a similar pattern to those of Ifi1 (Fig. 1c). This was expected as Stat1 regulates the transcription of both genes in the context of IFN-β activation. After an initial rapid increase of Ch25h RNA abundance, its RNA levels remained elevated between 4 and 10 h and subsequently decreased by 24 h. In these experiments, increases in Ch25h RNA abundance were not completely negated in the knockout BMDMs. This is likely due to the activation of IFNAR1 by other type I interferons [24]. Most notably, Atf3-induced levels were ablated in Poly(I:C)-treated Ifnb−/− BMDMs (Fig. 1d). This demonstrates Atf3 has a strict dependency on the IFN-β response for the transcriptional activation of its promoter upon TLIR stimulation. The molecular “wiring” for this inflammatory regulatory circuit is shown in Fig. 6, which details the stringent, coordinate control underpinning the regulation of 25-HC synthesis in IFN-activated BMDMs.

3.2. IFN-inducible-25-HC mediates a dose-dependent reduction of HMGCR protein levels in BMDMs

We next sought to quantitatively determine proteomic parameters for HMGCR inhibition that delineate the molecular circuitry for the IFN-inducible 25-HC regulation of the sterol pathway. Whilst the mechanism by which 25-HC regulates the SREBP pathway of the sterol pathway has been extensively characterised (summarised in Fig. 6), it is also known that 25-HC can post-translationally regulate HMGCR protein abundance in a SREBP independent manner [16,17]. We reasoned, therefore, that IFN-γ treatment might modulate the levels of Hmgcr mRNA and HMGCR protein through SREBP-dependent and independent mechanisms, respectively. To test this hypothesis, we quantified how increasing doses of 25-HC alter HMGCR protein abundance in BMDMs. Under normal serum conditions, endogenous HMGCR protein levels are
weakly detected when analysed by western blotting. HMGCR protein levels can, however, be markedly increased upon cholesterol depletion through compensatory homeostatic activity of the SREBP-activated pathway. We first quantified the HMGCR protein levels can, however, be markedly increased upon cholesterol depletion through compensatory homeostatic activity of the SREBP-activated pathway. We first quantified the HMGCR protein abundance in BMDMs quantitated following IFN-γ treatment in wild-type BMDMs (relative to control treated) in Medium A over an 8-h period. Each point represents transcript synthesis in control treated vs IFN-γ during a 30-min period. Normalised log fold change values were calculated by subtracting the control treated from the IFN-γ treated signal values. 

3.3. IFN-γ exerts an immediate early SREBP-independent proteomic layer of regulation of HMGCR followed by a delayed early SREBP-dependent transcriptional mechanism

In our next experiments, we aimed to determine and compare the extent of IFN induced reduction in HMGCR RNA levels (dependent on the activity of the SREBP pathway) and protein levels (which we assume involves a combination of SREBP-dependent and SREBP-independent pathways). Time course experiments measuring the reduction of total RNA of Hmgcr after 8 h of IFN treatment revealed RNA abundance for this transcript decreased approximately 50% reduction in RNA abundance by 8–9 h is...
due to changes in the rates of transcription alone and the determined turnover rates (approximately 73 min) suggest at least an hour response time to effect a change in abundance.

We next investigated the quantitative effects of direct 25-HC administration (2.5 µM) to BMDMs on RNA and protein levels under sterol-depleted conditions as discussed above. Consistent with the data in Fig. 3a and b, a significant reduction in RNA levels was detected at 9 h post-treatment revealing a greater magnitude of RNA reduction (approximately 45%) with only a small decrement at 1.5 h (Fig. 3d). In contrast, levels of HMGCR protein were more than 90% reduced by 1.5 h of 25-HC treatment and maintained a similar reduced level at 9-h post-treatment (Fig. 3e and f). In summary, these experiments show that 25-HC rapidly diminishes macrophage HMGCR protein levels within the first hour of treatment. Note, at the later time point (9 h), reductions in RNA levels will also contribute, in an additive manner, to a sustained reduction of HMGCR abundance.

We next quantified the RNA (Fig. 3g) and protein (Fig. 3h and i) levels of HMGCR with IFN-γ treatment at multiple time points. These experiments were also performed using sterol-depleted conditions. Fig. 3g–i shows that IFN-γ treatment results in a 20% decrease for both RNA and protein levels of HMGCR at 1.5 h post-treatment. However, at 9-h post-treatment, the reduction in RNA and protein levels is about 50% and 90%, respectively. These results point towards both transcriptional and post-translational mechanisms contributing to the maintenance of reduced HMGCR levels at late times. These data provide evidence showing that IFN-γ and 25-HC can transcriptionally and post-translationally alter levels of HMGCR. These differential modes of action are temporally coordinated with rapid early effects on HMGCR mediated at
the protein level and its subsequent sustained reductions involving equal contributions at the protein and RNA levels.

3.4. IFN-γ induces proteasomal degradation of HMGCR

It is well documented that 25-HC can promote the degradation of HMGCR protein through an ubiquitin-proteasome system [16,17]. We hypothesised, therefore, that IFN-γ, acting through an induction of 25-HC, can also promote the proteasomal degradation of HMGCR protein.

To test whether the reduction of HMGCR protein by IFN-γ in BMDMs involves proteasomal degradation, we utilised the proteasome specific inhibitor, MG132 [16]. The results shown in Fig. 4a and b demonstrate that MG132 treatment can inhibit the degradation of HMGCR induced by 25-HC as expected. Strikingly, 6 h of IFN-γ treatment in BMDMs also leads to a significant reduction...
in HMGCR levels (Fig. 4a and c) that is completely blocked by the proteasome inhibitor, MG132. We conclude from these experiments that 25-HC or IFN-γ treatment leads to the proteasomal degradation of HMGCR.

3.5. IFN-γ-elicited reduction of HMGCR protein absolutely requires the de novo synthesis of 25-HC

The experiments described above strongly suggest the possibility that IFN-γ alters levels of HMGCR through de novo synthesis of 25-HC. To unequivocally test this possibility, we utilised BMDMs derived from Ch25h−/− mice. The genetic ablation of Ch25h results in a complete inability of BMDMs from Ch25h−/− mice to produce 25-HC [26]. We anticipated, therefore, that IFN-γ would be unable to reduce HMGCR abundance in these knockout cells. Accordingly, Ch25h−/− and wild-type BMDMs were treated with 25-HC (as a positive control) or, IFN-γ and the levels of HMGCR protein were quantified (Fig. 5a–d). As expected, both 25-HC and IFN-γ treatment of wild-type BMDMs lead to a greater than 90% decrease in HMGCR protein levels by 9 h post treatment and which temporally reflects both SREBP dependent and independent pathways. However, at earlier times (4 h post-treatment), only a 50% reduction in HMGCR was observed for IFN-γ treatment in contrast to the nearly 100% reduction observed for 25-HC (Fig. 5a and b). This level of reduction can not be accounted for by the SREBP pathway alone because of the delineated 3–4 h lag period required for optimal induction of CH25H by IFN-γ and therefore, indicates that the 50% reduction at 4 h of IFN-γ is mainly due to the SREBP-independent pathway for targeted proteasomal degradation. In Ch25h−/− BMDMs, the reduction in levels of HMGCR by IFN-γ treatment at both 4 and 9 h were completely abrogated while 25-HC was capable in mediating a decrease of HMGCR protein in these cells at both time points (Fig. 5c and d). These experiments therefore provide direct evidence that the first 9 h of IFN-γ-activation reductions in HMGCR protein abundance require the de novo synthesis of 25-HC by Ch25h.

As the induction of CH25H is rapidly suppressed we anticipate that the effective time for 25-HC may be limited to the first 12–24 h. Yet we have shown that in IFN activated BMDMs the inhibition of the sterol pathway is maintained for at least a period of 48–72 h [24]. We therefore asked whether the synthesis of 25-HC is also exclusively required for longer-term suppression of HMGCR.

Fig. 5e shows the results of experiments measuring Hmgcr RNA levels in wild-type and Ch25h−/− BMDMs treated with IFN-γ or 25-HC for 24 h. In wild-type BMDMs, an expected 40-50% drop in RNA levels is detected for 25-HC treatment while it is notable for IFN-γ treatment, Hmgcr levels are further decreased by 60–70%. In Ch25h−/− BMDMs, Hmgcr RNA levels are significantly reduced but statistically higher than the wild-type levels. As a control, in Ch25h−/− BMDMs 25-HC treatment retains a similar level of inhibition of Hmgcr RNA. Therefore, these results demonstrate that while the early protein degradation of HMGCR is completely dependent

![Fig. 4. IFN-γ induces proteasomal degradation of HMGCR. (a) Wild-type BMDMs were pre-treated with MG132 in Medium C for 1 h and then treated with 25-HC or IFN-γ in the same culture medium for another 6 h. Western blot was performed to determine HMGCR protein levels. (b) Intensity values of HMGCR to tubulin. As MG132 has an effect on β-actin protein levels, tubulin was used as the internal control. Data are mean ± SEM (n = 4). *p < 0.05, determined with an unpaired Student’s t test. (c) Intensity values of HMGCR to tubulin. Data are mean ± SEM (n = 4). **p < 0.01, determined with an unpaired Student’s t test.](https://doi.org/10.1016/j.steroids.2015.07.023)
on the de novo synthesis of 25-HC, the longer-term suppression mediated at the RNA level is only partially (approximately 50%) dependent on this oxysterol.

3.6. A model for the regulatory control of HMGCR by IFN

The above experiments suggest a complex multi-layered regulatory pathway by which IFN-γ suppresses HMGCR. We have demonstrated that IFN-γ can transcriptionally and post-translationally regulate HMGCR, mediated through the de novo synthesis of 25-HC. Altogether, our results provide a foundation for further understanding of the complex regulation of the sterol biosynthesis pathway by the immune system through computational modelling. Toward this endeavour, Fig. 6 provides a graphical model for the coordinate regulation of HMGCR by IFN-γ, constructed using the systems biology graphical notation (SBGN) [54]. Specifically, IFN transcriptionally increases the expression of Ch25h via STAT1, whose translational product catalyses the production of 25-HC. The newly synthesised 25-HC can rapidly promote the proteasomal degradation of HMGCR and retain SREBP2 in the ER, which suppresses the transcription of Hmgcr and with concomitant temporal delay in translation further reduces HMGCR. The induction of Ch25h is curtailed by the repressor ATF3 that also negatively feedback on the IFN pathway.

4. Discussion

In this study we quantitatively elucidate how IFN-γ regulates the mevalonate-sterol pathway by targeting HMGCR through both SREBP-dependent and independent pathways. We show that in BMDMs, IFN-γ elicits a rapid proteasomal degradation of HMGCR. Importantly, the effects of this degradation are later potentiated through transcriptional and translational repression of HMGCR. Further, we demonstrate that IFN-γ mediates this effect through the de novo induction of CH25H expression and subsequent synthesis of 25-HC. This work provides, therefore, a foundational
framework for formal exploitation and advances new mechanistic insights into IFN-γ mediated macrophage suppression of mevalonate-sterol biosynthesis and the anti-viral response.

The mevalonate-sterol biosynthesis pathway is active in all cells of the body and is responsible for the production of cholesterol, isoprenoids, ubiquinone and downstream metabolites of oxysterols and steroid hormones. These products are involved in a range of core biological processes necessary for cell growth, metabolic and oxidative homeostasis, membrane integrity, vesicular transport and endocrine physiology. In addition, infectious agents, such as microbial, parasitic, and especially viruses, also rely on biosynthetic cellular pathways for their growth [55]. A common feature of many unrelated pathogens is a dependency on cellular lipid metabolism and we, and others, have shown that pharmacological inhibition of the mevalonate-sterol biosynthesis pathway, in particular, can curtail pathogen growth [32,33,24,34]. Increasingly, studies are emerging that the immune system and lipid pathways are tightly coupled. Notably, both upstream and downstream metabolites of this pathway are known to play key roles in immunity and likely explain its long standing association with inflammatory based diseases such as atherosclerosis. Defining how immunity and lipid metabolism are integrated, share resources and cross-regulate one-another during infection is, therefore, of great importance to our understanding of anti-infective immunity and the development of new therapeutic strategies.

Recently, we identified an interferon-induced reduction in sterol biosynthesis as a key component of the very early cellular response to virus-infection in BMDMs [24]. This led to a focus on characterising the close coupling between interferon and mevalonate-sterol metabolism and to the identification of 25-HC biosynthesis as an important component of the interferon-induced anti-viral response in BMDMs [42,24,31]. For over 40 years, 25-HC was recognised to have a potent ability to negatively feedback on sterol metabolism and to the identification of 25-HC biosynthesis as an important component of the interferon-induced anti-viral response in BMDMs [42,24,31]. For over 40 years, 25-HC was recognised to have a potent ability to negatively feedback on sterol metabolism and to the identification of 25-HC biosynthesis as an important component of the interferon-induced anti-viral response in BMDMs [42,24,31].

Here, in our detailed kinetic and quantitative investigations, we systematically involve a temporal, multi-layered suppression of the mevalonate arm of the sterol biosynthesis pathway. Within the first 90–200 min of IFN-γ stimulation of BMDMs, HMGCR is directed by SREBP-dependent and independent mechanisms, 25-HC can inhibit, in a cell-autonomous manner, a wide range of viruses. The principal pathway for anti-viral activity mediated by 25-HC for a number of viruses consistently maps to the proximal mevalonate arm of the pathway. Importantly, HMGCR represents the first regulatory enzymatic step in this part of the sterol pathway and statins or interference RNA inhibition of HMGCR markedly inhibit viral growth. The role of 25-HC anti-viral pathway in vivo requires further investigation.

How precisely IFN controls the sterol pathway is little understood. Here, in our detailed kinetic and quantitative investigations, we further explore the IFN regulation of sterol biosynthesis, we systematically uncovered evidence that demonstrates a direct coupling between IFN-γ signalling and the synthesis of 25-HC that mechanistically involves a temporal, multi-layered suppression of the mevalonate arm of the sterol biosynthesis pathway. Within the first 90–200 min of IFN-γ stimulation of BMDMs, HMGCR is directed by 25-HC for proteasomal degradation and is coordinately sustained for reduced expression through blockade of the restorative SREBP pathway, suppressing both RNA and protein synthesis (Fig. 6). Thus, our findings provide a foundation and framework for future modelling and predictive therapeutic intervention strategies for regulating sterol biosynthesis and its immune effector functions.

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