Cell fusion in the liver, revisited

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Abstract

There is wide agreement that cell fusion is a physiological process in cells in mammalian bone, muscle and placenta. In other organs, such as the cerebellum, cell fusion is controversial. The liver contains a considerable number of polyploid cells: They are commonly believed to originate by genome endoreplication, although the contribution of cell fusion to polyploidization has not been excluded. Here, we address the topic of cell fusion in the liver from a historical point of view. We discuss experimental evidence clearly supporting the hypothesis that cell fusion occurs in the liver, specifically when bone marrow cells were injected into mice and shown to rescue genetic hepatic degenerative defects. Those experiments—carried out in the latter half of the last century—were initially interpreted to show “transdifferentiation”, but are now believed to demonstrate fusion between donor macrophages and host hepatocytes, raising the possibility that physiologically polyploid cells, such as hepatocytes, could originate, at least partially, through homotypic cell fusion. In support of the homotypic cell fusion hypothesis, we present new data generated using a chimera-based model, a much simpler model than those previously used. Cell fusion as a road to polyploidization in the liver has not been extensively investigated, and its contribution to a variety of conditions, such as viral
About 70% of hepatocytes are polyploid, arising either from genome duplication without division (endoreplication) or from cell fusion. Experiments with chimeric mice containing two cell populations each bearing a different genetic marker had shown that some liver cells express markers of both genomes, suggesting that cell fusion occurred. Here, we review the data in the literature and describe new experiments using a chimeric model that confirms that cell fusion contributes to liver polyploidy. We argue that the role of cell fusion in pathological conditions, such as viral hepatitis and neoplastic transformation, is worth further study.

INTRODUCTION
Mammalian cells are usually diploid, with the exception of mature gametes, which are haploid. Interestingly, a few tissues contain polyploid cells, such as muscle cells, osteoclasts, hepatocytes, megakaryocytes and trophoblasts[1]. A common feature in all these tissues is the presence of a diploid progenitor cell that at some point during the differentiation/maturation process becomes polyploid. Polyploidization can be explained by two main mechanisms: endoreplication and cell fusion. Endoreplication occurs when the genome is duplicated without cell division, whereas cell fusion occurs between two different cells, either of the same or of a different identity. In the latter case, the genomes from two different cell types—which can come even from different species—fuse within the same membrane and, therefore, coexist within the same cell[2]. For both mechanisms, the outcome is a polyploid cell.

POLYPLOIDY IN NORMAL CELLS
It is generally recognized that polyploidization in normal organisms is adaptive since it helps specialized cells acquire the ability to perform new, specific functions[3]. For example, osteoclasts are large multinucleated cells that perform the difficult task of resorbing bone matrix. This specialized function cannot be accomplished, for example, by mononucleated TRAP+ osteoclasts as this leads to osteopetrosis, a disease in which bone is not degraded[4]. Among the other types of polyploid cells, there is agreement that muscle cells and trophoblasts are products of cell fusion. In contrast, megakaryocytes are polyploid cells generated by genome duplication followed by aborted cytokinesis[5].

Polyploid liver cells are usually considered to be formed by endoreplication of the genome[6-7]. This conclusion was originally based on the seminal work of Mintz and colleagues, who pioneered the use of chimeric mice to study gene expression[8]. In the chimeric mice, cells with two different genomes coexist in a single organism. If a cell fuses with another having a different genome, the resultant cell will contain markers of both. In their studies, Mintz and colleagues concluded that fusion occurred in muscle but not in the liver[8-10]. However, they had to rely mainly on the analysis of isoforms expressed by tissue-specific enzymes because single-cell markers were not yet available. Contrary to osteoclasts, trophoblasts and muscle cells, whose nuclei maintain their individuality, polyploid liver cells can be bi- or mono-nuclear.

Specific cell cycle genes, such as Cdk1, are involved in polyploidy formation and maintenance in liver[11]. Interestingly, after partial hepatectomy, quiescent hepatocytes can start proliferating again: deletion of cyclin-dependent kinase 1 (Cdk1), 2 (Cdk2), cyclin E1 or E2 individual genes does not limit liver regeneration, but concomitant ablation of Cdk2 and Cyclin E1 reduces liver regeneration, suggesting partial overlapping function of some cell cycle genes[11,12].

TRANSDIFFERENTIATION AND FUSION:
A DEBATE NOT YET SETTLED
The study of cell fusion in the liver began while investigating the existence of lineage transdifferentiation. The possibility of cell transdifferentiation was raised at the end of the last century in a paper published in Science[13] in which the authors, after having transplanted neural stem cells transgenic for the beta-galactosidase (βgal) gene into wild-type mice, detected βgal+ cells in peripheral blood. Their interpretation was that neural cells had transdifferentiated into cells of the hematological lineage. This plasticity was surprising to many but, because the report was published just after the birth of Dolly the sheep, looked plausible[14]. We must emphasize here that that work had nothing to do with the reprogramming approach reported several years later by Yamanaka, who, in contrast, obtained reprogramming by forced expression of intracellular transcription factors[15]. The plasticity of neuronal stem cells was claimed by Bjornson et al[13] to occur by simple exposure to endogenous factors present in vivo.

Essentially, studies on plasticity were performed by transplanting cells from a mouse transgenic for an easily detectable marker gene (e.g., βgal) into a non-
transgenic animal. The appearance of βgal-marked cells in an organ different from the tissue of origin was interpreted to be the result of transdifferentiation. The Science paper in which cells of the nervous system were suggested to acquire a hematological fate, was rapidly followed by other examples of transdifferentiation involving cells of several other lineages, including blood, brain, muscle, kidney and heart.[16-25].

Due to the high potential for translation to the clinic, bone marrow cells (BMCs) escalated to center stage. BMCs are easily obtained, extensively investigated, routinely transplanted and well-characterized in humans. If simple transplantation protocols allowed the rescue of degenerative defects in organs such as brain, kidney or liver, we would have a sort of panacea in hand. Unfortunately, although many clinics-mostly in the United States-still advertise these kinds of treatments[26], transdifferentiation as originally proposed in the Science paper has not been confirmed by subsequent, more controlled studies[27-32]. Indeed, although a limited transdifferentiation capacity of some cells cannot be completely ruled out, more-recent studies have shown that transdifferentiation is often an experimental artifact. As stated above, transdifferentiation was claimed to occur if, after a given lineage (for example, hematopoietic cells) expressing a reporter gene was transplanted into a wild type mouse, cells of other lineages (for example, brain) were found to coexpress the reporter gene with accepted markers of their lineage. While βgal was initially used as a marker, most subsequent papers exploited fluorescent reporter genes that could be easily traced in vivo. It was assumed that all fluorescent cells found in a normal, non-transgenic, mouse had to be the progeny of transgenic donor cells: Hence, if they were found in other organs, they must have derived from original cells that had acquired a new fate by transdifferentiation.

In addition to trivial technical artifacts, cell fusion was raised to explain some of these results: in the experimental design discussed above, fusion between any cells of the host with transplanted donor cells could have provided the former with the reporter gene. It is difficult to discriminate between the two possibilities—transdifferentiation and cell fusion—with simple marker analysis.

As mentioned above, transdifferentiation of BMCs would be an attractive approach for regenerative medicine. Heart, brain and liver are heavily affected by degenerative genetic diseases that have a huge impact on human health; they would all greatly benefit from cell fusion-based therapies using exogenous cells, if that mechanism indeed occurs in vivo. Certainly, exogenous cells could provide defective endogenous ones with the missing genetic component while maintaining the differentiation status of the mature cell.

With regard to the liver, several reports in which BMCs were transplanted into recipient mice in the hope of inducing hepatocyte transdifferentiation showed that cells bearing donor-derived cellular markers could be found in host livers[18-21]. These “transdifferentiated” cells increased in number when the host livers were either injured (partial hepatectomy) or affected by a chronic degenerative genetic defect. However, the results were challenged by scientists who were unable to reproduce the transdifferentiation of hematological cells into non-hematological ones[33-36]. Cell fusion was shown to occur in vivo, so several reports investigated fusion events in a variety of other models. In the liver, the most spectacular experiments were performed by Grompe’s group on the classical model of fumarylacetoacetate hydrolase (Fah) deficiency[37]. Mice recessive for a Fah mutation are models for tyrosinemia type I, a severe genetic disease leading to liver failure in humans. Grompe and coworkers showed that bone marrow transplants in these mice led to the generation of liver cells bearing the donor marker, and demonstrated that this event was not due to transdifferentiation of hematological into hepatic lineage cells. Instead, these marker-carrier liver cells originated from cell fusion between donor bone marrow and resident hepatocytes, leading to polyploid cells that were not easily distinguishable from true hepatocytes in that the latter could also be polyploid. Due to the growth advantage shown by normal hepatocytes over diseased ones, the approach was so efficient that several mice were essentially cured. Results were confirmed by further studies[38-40], which also pointed to macrophages as the hematological cell responsible for fusion[41,42]. These results are in agreement with macrophages being physiologically prone to cell fusion[43]. In a review of 77 published studies on the generation of hepatocytes by hematopoietic cells transplanted in liver, the authors concluded that cell fusion was the mechanism involved[44]. Cell fusion is enhanced by the presence of liver injury or chronic disease, such as in the Fah model, since in a well-controlled study in which BMCs were injected into normal recipients, only 7 out of 470000 liver cells examined bore donor markers as a result of cell fusion[45]. In addition to BMCs, other types of cells, such as mesenchymal or amniotic stem cells and cells differentiated from pluripotent stem cells, can fuse with cells in injured livers, even when injected into a different species[45]. Human umbilical cord blood cells have also been reported to fuse with hepatocytes of immunocompromised mice[46], although no evidence of cell fusion was reported in other studies[47-49]. Moreover, cell fusion and transdifferentiation have been claimed to coexist[50].

The cell fusion-based explanation was found to hold also in other similar experimental settings[34,37,39,51,52] (reviewed in[27,53,54]). However, the possibility that at least in some cases, especially when an injury is applied to the recipient organ, bone marrow donor cells could be directed toward a different fate has not been completely ruled out, since several reports of well controlled differentiation have been published[55-64].
CELL FUSION IN THE NORMAL LIVER

The discovery that cell fusion can cure a degenerative disease of the liver prompted Grompe’s group to investigate whether cell fusion occurs also in the disease-free state. The experimental plan to address this was as follows: they transplanted $1 \times 10^5$ wild-type (Fah$^{-/-}$) hepatocytes into each of four Fah$^{-/-}$/βgal$^+$ recipients. After more than 80% of the liver was repopulated, $1 \times 10^5$ hepatocytes were serially transplanted into each of two Fah$^{-/-}$ recipients and the liver was again repopulated to a donor contribution of more than 80%. Then they analyzed $3 \times 10^7$ Fah$^+$ hepatocytes, but were unable to find a single Fah$^+$/βgal$^+$ cell. They concluded that the frequency of cell fusion, if any, was very low$^{[41]}$.

It must be taken into considerations that the protocol involved damaged livers and injections of adult cells. However, although complex, the approach looks suitable to address the question of cell fusion in the disease-free liver. The only caveat is that, if the originally transplanted wild-type cells were mature hepatocytes (the age of the mice used was not specified), then it is possible that they represent polyploid cells that were already fully differentiated and functional and, therefore, less prone to fuse. This is because at this stage they have already achieved the benefits of being large cells with multiple genomes.

Apart from the original studies by Mintz and colleagues already cited, other studies investigating whether cell fusion occurs in the normal liver are lacking. Cell fusion has occasionally been reported to occur in hepatocytes or in hepatic tumor lines cultured in vitro$^{[65-67]}$. Yet, this does not prove that the mechanism is physiologically relevant in vivo. For these reasons, Faggioli and coworkers devised and implemented a relatively simple but straightforward protocol based on chimeric mice, as originally proposed by Mintz’s group$^{[8-10]}$. Embryo-derived mouse chimeras are mice born from embryonal cells carrying different genomes$^{[49]}$. They can be created either by morula aggregation or by injection of embryonic stem cells (ESCs) into blastocysts, and they can be exploited for the study of cell fusion. If each of the two aggregated morulae contains a different reporter gene, then cells positive for both reporters will definitively be fused cells.

Faggioli et al.$^{[71]}$ reasoned that by aggregating morulae from two different strains of transgenic mice expressing either green fluorescent protein (GFP)$^{[69]}$ or the βgal protein (Rosa 26 mouse$^{[50]}$), the outcome would be animals that display two genetically distinct liver cell populations, each bearing a single marker (either GFP or βgal); any cell displaying both markers must be the result of cell fusion. With the appropriate controls, they identified three populations: GFP$^+$/βgal$^+$; GFP$^+$/βgal$^-$; and GFP$^-$/βgal$^+$. The percentage of double-positive cells in the chimeric samples was estimated to be about 25%.

The authors confirmed their results with two other independent strategies. Briefly, they performed PCR amplification on single hepatocytes with primers specific for each reporter gene, finding cells displaying both markers only in chimeric mice, in a percentage close to 10% of cells bearing at least one marker. In addition, the authors used fluorescent in situ hybridization (FISH) to investigate the sex chromosome content of hepatocytes in XY↔XX chimeric mice. They reasoned that, if fusion occurred between a female and a male cell, some binucleated cells containing Y chromosome(s) only in one of the two nuclei would be detected. Similarly, if mononucleated polyploid hepatocytes were analyzed, they should contain only X chromosomes in various numbers in the case they derived from the X component of the chimera mouse, or as many X as Y chromosomes if derived from an XY cell. In contrast, if the mononucleated polyploid hepatocytes were products of a cell fusion event between a female and a male cell, an unbalanced complement of X and Y chromosomes would be found. In the end, sex chromosome patterns were detected that were clearly indicative of cell fusion in binucleated as well as mononucleated hepatocytes in about 5%-10% of cells$^{[71]}$.

These results are at odds with those presented by Willenbring et al.$^{[41]}$. This discrepancy could be explained by the different approaches used, since that of Faggioli et al.$^{[71]}$ mimics normal liver development, while the one used by Willenbring et al.$^{[41]}$ involves the injection of exogenous hepatocytes into damaged liver and complex transplantation experiments. As mentioned before, this could ultimately lead to underestimation of fusion events in the latter study.

We are not aware of recent studies investigating cell fusion in normal liver, although replication of the chimera studies would not be too time consuming. Apparently, fusion is neither considered to occur frequently nor to be of physiological relevance. For this reason, while performing a study on the role of cell fusion in cancer$^{[72]}$, we addressed cell fusion with an even simpler approach based on the production of chimeric Cre: tdTomato mice. Morulae derived from mice transgenic for Cre recombinase under the control of a constitutive promoter were fused to morulae from mice transgenic for an inactive floxable tdTomato gene that is activated only if Cre recombinase is expressed in the same cell (Figure 1A). Cells from the two morulae will develop independently and no cell will be tdTomato-positive unless fusion with a Cre-containing cell has occurred (see the schematic representation in Figure 1B).

This approach—which has been widely used for lineage and transplantation studies—has the advantages of having an undetectable background if cell fusion does not occur, no interference between the two fluorescent reporter genes, and simple assessment in liver sections with well-validated tdTomato-specific antibodies. In addition, leakiness of the promoter, which sometimes occurs in Cre-based conditional mice, does not affect this model.
Analysis performed to date on two chimeric mice has clearly identified the presence of tdTomato-positive cells in the liver (Figure 2). Positivity was not detected in wild-type mice or in inactive tdTomato mice. As expected, the progeny of tdTomato-Cre mice crosses were positive in all tissues.

Fused cells are distributed all over the liver parenchyma, but are often found in clusters. This is in keeping with cell fusion occurring in cells maintaining their proliferative capacity, giving rise to a progeny that expands but remains in close proximity to their original location. This is in agreement with other studies showing hepatocytes originating from clonally derived clusters in postnatal liver\[73,74\]. However, the devil is in the detail and we are always at risk of artifacts\[75\]. In the chimeric experimental design, coexpression in the same cell of two reporter genes originally expressed independently by two distinct cells is commonly accepted as proof of a fusion event. This assumption was used in our original work on cell fusion\[71\]. However, the detection of fluorescence is prone to artifacts caused by endogenous background fluorescence, a phenomenon especially marked in liver; in the case of fIgA, endogenous enzymatic activity can also lead to misinterpretation. In addition, it has become increasingly appreciated over the last ten years that transfer of materials-including RNA and proteins-between cells via extracellular vesicles is a frequent phenomenon\[76-78\]. Therefore, it cannot be excluded that in the Cre-tdTomato approach aforementioned, RNA encoding Cre recombinase or tdTomato could have been transferred from the Cre\[+\]cell to the tdTomato one, and thus activating the reporter locus leading to expression of the reporter protein. Even the transfer of a few RNA or protein molecules over a very short period of time can activate the tdTomato gene, which then would become permanently expressed. However, the Cre-Lox and GFP systems have been widely used, in general giving consistent results for expression and expected specificity. Unfortunately, with the technologies available to date there is no way of discriminating fusion events from vesicle-mediated transfer in vivo while maintaining physiological conditions. In this regard, it is worth mentioning that several recent papers analyzing the fate of GFP\[+\]cells transplanted into mouse retina have reported the detection of GFP\[+\]cells that did not originate from the donor\[79-81\]. This suggests that GFP activity was leaked into the intracellular space and absorbed by endogenous cells or was transferred to them by extracellular vesicles—fusion can be excluded since retinal cells were normal in size and not polyploid. This is troubling if true, and some lineage
or transplantation studies based on the detection of reporter genes should be carefully re-examined. Techniques based on in situ hybridization with probes specific for sex chromosomes can be used to demonstrate cell fusion \[71\], since the presence of an XY nucleus as well as an XX one in a binucleated cell should definitively be due to cell fusion. This technique— which does not allow the analysis of live cells—has been used in studies on the ploidy of hepatocytes, with the caveat that the analysis might be complicated by the aneuploidy shown by some normal human and murine liver cells \[82-85\]. In any case, it will be difficult to investigate cell fusion in man: in theory, transplantation of male hepatocytes in female hosts performed for regenerative liver diseases could detect cell fusion, but this is a very rare occurrence and would require biopsies or post-mortem examination.

CONCLUSION

Cell fusion in the liver is still controversial. Thus, replication of previous studies with appropriate mouse chimeras is welcomed. Endoreplication and cell fusion are not mutually exclusive, as suggested by Gentric and Desdouets \[86\]. We strongly believe that fusion in the liver should be studied in order to confirm and explain this phenomenon. If established, this will open several new lines of investigation. For example, is cell fusion or endoreplication preferred in different contexts, or are they interchangeable? What is the fusion potential of hepatocytes with a DNA content higher than 4n? Are there hepatocytes with unbalanced or uneven-n chromosome numbers, and are there fusion products between one diploid and one tetraploid cell? Does cell fusion occur in species other than rodents, and particularly in man? Can fused cells participate in the ploidy reduction occurring after partial hepatectomy? Are HBV or HCV infections, which are themselves fusogenic viruses, able to change hepatocyte ploidy and binuclearity \[87\], or do other metabolic stresses \[88\] affect endoreplication or fusion? Does cell fusion play a role in HCV-mediated liver carcinogenesis \[89\]?

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