

1 **Ferroptosis and metabolic dysfunction-associated fatty liver**
2 **disease: is there a link?**

3 **Short Title:** Ferroptosis in MAFLD

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36 **Abbreviation list**

37 GPX4 = glutathione peroxidase, GSH = glutathione, 4-HNE = 4-hydroxynonenal,
38 HSC = hepatic stellate cells, IR = Insulin resistance, MAFLD = metabolic
39 dysfunction-associated fatty liver disease, NAFL = nonalcoholic fatty liver, NAFLD =
40 non-alcoholic fatty liver disease, NAS = NAFLD activity score, NASH = non-
41 alcoholic steatohepatitis, OR = odds ratio, PDFF = proton density fat fraction, ROS =
42 reactive oxygen species, T2DM = type 2 diabetes mellitus

43

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55 All authors have nothing to declare.

56

57 **Authors' contributions**

58 Gong Feng and Ming-Hua Zheng designed the review outline. Gong Feng collected,
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63

64 **Data availability statement**

65 Some or all data, during the review are available from the corresponding author by
66 request.

67

68 **Abstract**

69 Non-alcoholic fatty liver disease (NAFLD), recently re-defined and re-classified as
70 metabolic dysfunction-associated fatty liver disease (MAFLD), has become
71 increasingly prevalent and emerged as a public health problem worldwide. To date,
72 the precise pathogenic mechanisms underpinning MAFLD are not entirely
73 understood, and there is no effective pharmacological therapy for NAFLD/MAFLD.
74 As a newly discovered form of iron-dependent programmed cell death, ferroptosis can
75 be involved in the development and progression of various chronic diseases, but the
76 pathogenic connections and mechanisms that link MAFLD and ferroptosis have not
77 been fully elucidated. The main characteristics of ferroptosis are the accumulation of
78 lipid peroxides and reactive oxygen species. In this brief narrative review, the
79 mechanisms of ferroptosis and its putative pathogenic role in MAFLD are discussed
80 to highlight potential new research directions and ideas for the prevention and
81 treatment of MAFLD.

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83 **Key-words:** ferroptosis; MAFLD; pathogenesis, type 2 diabetes, metabolic
84 syndrome

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95 **Introduction**

96 In the past, two main mechanisms of programmed cell death have been recognized,
97 namely apoptosis and necrosis. However, these two mechanisms do not explain all
98 forms of cell death. Other forms of programmed cell death have recently been
99 recognized, such as pyroptosis, autophagy and ferroptosis. Of these newly discovered
100 forms of programmed cell death, ferroptosis has been widely studied in recent years,
101 because it involves accumulating lipid-active oxygen species and appears to play a
102 key role in the development of some types of tumors, neurodegenerative diseases, or
103 endocrine diseases.¹⁻³

104

105 Non-alcoholic fatty liver disease (NAFLD), recently named as metabolic dysfunction-
106 associated fatty liver disease (MAFLD)^{4,5,6}, has reached epidemic proportions,
107 becoming the most common cause of chronic liver diseases worldwide (affecting up
108 to ~30% of world's adults).^{7,8} Growing evidence supports the notion that MAFLD is a
109 “multisystem” disease, in addition to causing severe liver damage (i.e., MAFLD-
110 related cirrhosis)⁹, affecting the vasculature and other organ systems that requires a
111 multidisciplinary and holistic approach.¹⁰

112

113 To date, the role of ferroptosis in the development and progression of MAFLD has not
114 been fully elucidated. Therefore, in this review we briefly discuss the role of
115 ferroptosis in MAFLD in order to highlight new areas for potential research into the
116 prevention and treatment of this common and burdensome liver disease.

117

118 **1. Overview of ferroptosis**

119 *1.1 The concept and characteristics of ferroptosis*

120 In 2012, Dixon et al. first coined the term ferroptosis as a new form of regulated cell
121 death.¹¹ This form of cell death results from glutathione depletion and glutathione
122 peroxidase inactivation (**Figure 1**).^{10,12} As an iron-dependent form of non-apoptotic
123 regulated cell death, iron plays a key role in the occurrence of ferroptosis.

124 Experimentally, it has been shown that iron-chelating agents may inhibit ferroptosis.¹³

125 At the same time, imbalance of iron metabolism induces lipid peroxidation and
126 production of reactive oxygen species (ROS), thereby triggering ferroptosis. The
127 regulatory mechanisms of ferroptosis are closely related to reactive oxygen clusters,
128 and accumulation of reactive oxygen clusters triggered by the Fenton-like reaction,
129 nicotinamide adenine dinucleotide phosphate-dependent lipid peroxidation, and
130 glutathione depletion.¹ Thus, as an atypical oxidative form of regulated cell death,
131 ferroptosis may induce cell death by increasing ROS production, thereby affecting
132 development of disease.

133

134 *1.2 Comparison of ferroptosis with other forms of programmed cell death*

135 From a morphological point of view, apoptosis is characterized by the occurrence of
136 typical apoptotic cellular bodies and by no rupture of cell membranes.¹⁴ Cell necrosis
137 is characterized by cell swelling, nucleus concentration, fragmentation, and
138 dissolution, as well as chromatin staining and flocculent, and organelle enlargement or
139 fragmentation. In contrast, ferroptosis does not show any typical morphological
140 features of both apoptosis and necrosis. Ferroptosis is typically characterized by cell
141 membrane rupture and vesiculation, reduced mitochondrial cristae, mitochondrial
142 atrophy, as well as lack of chromatin agglutination in the nucleus.¹⁴

143

144 In terms of biochemical characteristics and regulatory mechanisms, apoptosis is
145 mainly dependent on cysteinyl aspartate-specific proteinase. During cell apoptosis,
146 Ca^{2+} and pH levels of cytoplasm increase, and endonuclease is activated, leading to
147 nuclear DNA fragmentation. Cell necrosis induces a severe local inflammatory
148 response that is associated with activation of signaling pathways, such as receptor-
149 interacting protein kinase 3. In contrast, in ferroptosis, intracellular Fe^{2+} accumulation
150 occurs, and levels of lipid peroxidation increase significantly.¹⁵ At the same time,
151 ROS production increases, cellular cystine/cysteine uptake decreases, glutathione
152 (GSH) is reduced, and some mediating factors, such as arachidonic acid, are
153 released.¹⁵ The essential nature of ferroptosis is intracellular Fe^{2+} accumulation that is
154 a typical disorder of cell redox metabolism. Thus, intracellular antioxidant capacity

155 decreases, and ROS and lipid peroxidation products accumulate in large quantities,
156 inducing cell death. There is no overlap between the regulatory mechanisms of
157 ferroptosis and those implicated in cell apoptosis or necrosis, and small molecules that
158 inhibit cell apoptosis and necrosis do not have any inhibitory effect on ferroptosis.¹⁶

159

160 **2. Effects of regulatory mechanisms of ferroptosis on MAFLD**

161 MAFLD comprises a histological spectrum of progressive liver conditions, ranging
162 from simple steatosis to steatohepatitis, advanced fibrosis and cirrhosis. The
163 pathophysiology underlying MAFLD involves a multitude of interlinked processes,
164 including insulin resistance (IR), lipotoxicity attributable to the accumulation of toxic
165 lipid species, infiltration of proinflammatory cells causing hepatic injury and
166 ultimately leading to hepatic stellate cell (HSC) activation and increased liver
167 fibrogenesis.

168

169 ***2.1 Lipid peroxidation***

170 Ferroptosis was first detected after stimulation of RAS-mutant cells by a small
171 molecular substance (i.e., erastin).¹⁷ Subsequent studies have shown that lipid
172 peroxidation is one of the major drivers of ferroptosis.¹⁸ Lipids are widely found in
173 biofilms and lipoproteins. When lipid peroxidation occurs, lipids become the major
174 target of peroxidation by increasing ROS. In turn, the increased production of ROS
175 leads to increased oxidative stress, and oxidative stress-induced lipid peroxidation
176 could play a key role in the development and progression of MAFLD.¹⁹ Convincing
177 evidence supports the importance of different lipid peroxidation products in the
178 development of MAFLD. Among the different aldehydes that can be formed as
179 secondary products during lipid peroxidation, malondialdehyde (MDA) and 4-
180 hydroxynonenal (4-HNE) are the two most extensively studied and both are
181 associated with different stages of MAFLD. A significant association between hepatic
182 4-HNE adducts and increasing stage of fibrosis has been described, and increased
183 mitochondrial 4-HNE–protein adducts during MAFLD development have been

184 reported.^{20,21} Studies also reported higher circulating levels of MDA, as measured by
185 the 2-thiobarbituric acid reaction assay, and higher levels of low-density lipoprotein
186 (LDL) oxidation in patients with MAFLD than in control subjects.^{22,23}

187

188 **2.2 Iron overload**

189 Not only is abnormal lipid metabolism involved in the development of MAFLD, but
190 also imbalance of iron metabolism may affect the occurrence of MAFLD.²⁴ Because
191 iron has two different valence states, iron participates in intracellular redox reactions
192 *in vivo*, enabling iron to produce oxidative free radicals. Iron overload induces
193 oxidative stress by producing ROS. The liver is one of the most critical organs for
194 iron storage. Approximately 25-30% of total iron in the body is stored in ferritin in the
195 liver, and the intrahepatic contents of iron and ROS are greater in diseased liver than
196 in normal liver, suggesting that ferroptosis may be associated with chronic liver
197 diseases.²⁵

198 A cross-sectional study, involving 5445 Chinese individuals, showed that there was a
199 dose-response relationship between dietary iron intake and the prevalence of
200 MAFLD.²⁶ In a case-control study in Southeast China, Pan et al. also found that
201 elevated serum ferritin levels were associated with a higher risk of MAFLD (adjusted-
202 odds ratio 1.62, 95% CI 1.16-2.27), and the hepcidin-to-ferritin ratio was significantly
203 associated with a lower risk of MAFLD (adjusted-odds ratio 0.70, 95% CI 0.50-0.98).
204 ²⁷ In a proof-of-concept study, Rostoker et al. prospectively analyzed the association
205 between intra-hepatic iron content and magnetic resonance imaging-proton density fat
206 fraction (MRI-PDFF) in 68 patients on chronic dialysis.²⁸ Among these dialysis
207 patients, 17 patients were followed-up during the period of iron therapy. The results of
208 this pilot study showed that liver fat content (assessed by MRI-PDFF) of patients with
209 moderate or severe iron overload was higher than that of normal iron load patients or
210 mild iron overload patients [median (interquartile range) MRI-PDFF-assessed liver fat
211 content: 7.9% (0.5-14.8%) vs. 5.0% (0.27-11%) vs. 5.0% (0.30-11.6%), respectively,
212 $P < 0.05$]. In 7 patients who received iron treatment, both liver iron and fat contents

213 increased concomitantly, whereas in 10 patients with iron overload, liver fat content
214 decreased after parenteral iron withdrawal, thereby suggesting that liver iron load may
215 influence liver fat content in these dialysis patients.²⁸ Barrera et al. also demonstrated
216 that in rats, iron overload significantly increased hepatic fat content, serum
217 transaminase levels, and induced a disruption in the desaturation capacity leading to
218 polyunsaturated fatty acid (PUFA) depletion, all of which were diminished by
219 antioxidant intervention.²⁹ Wang et al. showed that high iron levels served as a driving
220 factor in the induction of ferroptosis, and the ferroptosis could damage liver
221 mitochondria associated with elevated serum ALT levels.³⁰ It is known that
222 inflammation and fibrosis are two critical stages in the pathophysiology of MAFLD.
223 Meanwhile, iron is known to increase the respiratory burst activity of Kupffer cells,
224 which may have a proinflammatory impact through the activation of nuclear factor
225 (NF)- κ B, thereby triggering the hepatic production of multiple pro-inflammatory and
226 fibrogenic mediators.³¹ Finally, in a small intervention study, Yamamoto et al. showed
227 that dietary restriction of calories, fat and iron improved the grade of hepatic iron
228 accumulation and oxidative stress in patients with MAFLD. In addition, the levels of
229 serum ALT and ferritin were significantly decreased.³²

230 It is known that IR is a pathogenic factor in the development and progression of
231 MAFLD.³³ Previous studies found that excessive iron accumulation may adversely
232 affect insulin secretion from pancreatic β -cells and, at the same time, may interfere
233 with expression of insulin receptors, thus resulting in greater IR. Pancreatic β -cells are
234 highly sensitive to levels of iron ions and can express hepcidin, which relieves iron
235 overload.³⁴ Moreover, excessive iron accumulation also induces oxidative stress and
236 mitochondrial damage, thereby further impairing pancreatic β -cell function. Hepatic
237 iron overload may also exacerbate hepatic IR by directly damaging liver cells.^{34,35}

239 **2.3 Glutathione (GSH)**

240 As discussed above, ferroptosis is a modulated form of cell death that is characterized
241 by the iron-dependent accumulation of lipid peroxidation to lethal levels. When

242 cystine transport proteins are inhibited (e.g., erastin), intracellular GSH is depleted,
243 resulting in inactivation of glutathione peroxidase (GPX4) and accumulation of lipid
244 peroxidation products that induce cell death.^{36,37} Koruk et al. showed that serum GSH
245 levels were higher in patients with MAFLD than in controls, suggesting that GSH
246 might play a key role in MAFLD pathogenesis and disease progression.³⁸

247

248 **3. Impact of ferroptosis on MAFLD**

249 ***3.1 Impact of ferroptosis on MAFLD-related risk factors***

250 ***3.1.1 Obesity***

251 Previous studies have shown that MAFLD is closely related to obesity, and ferroptosis
252 might further promote the development of MAFLD by affecting obesity.^{39,40}

253 Experimentally, it has been reported that iron-chelating agents (e.g. deferoxamine,
254 deferiprone or deferasirox) have the potential to treat obesity. For instance, Yan et al.
255 reported that deferoxamine could reduce the expression of fat-generating genes in
256 adipose tissue and reduce the expression of genes related to mitochondrial
257 biosynthesis, thereby achieving the effect of treating obesity.⁴¹ In another
258 experimental model of polygenic obese mice, Ma et al. found that increased iron
259 concentration is associated with adipose tissue remodeling and increased adipose
260 tissue IR.⁴² However, further experimental research is required in this field.

261

262 ***3.1.2 Type 2 diabetes***

263 Apart from well-studied IR in peripheral cells, impaired insulin secretion from
264 pancreatic β cells has been acknowledged as the core defect in the development of
265 T2DM, because of exhaustion of pancreatic β cells and consequent failure of insulin
266 secretion.⁴³ Pancreatic β cell death, involving the modulation of both pancreatic β cell
267 mass and function, is believed to be the final pathogenic event in the progression of
268 T2DM, leading to rapid deterioration of glycaemic control, if not treated properly.⁴⁴
269 Li et al. reported that ferroptosis may contribute to pancreatic β cell loss and
270 dysfunction, and insulin secretion is worsened by ferroptosis-inducing erastin or
271 RAS-selective lethal compounds.⁴⁵ Quercetin is a potential glucose-lowering

272 supplement that might also exert some beneficial effects on ferroptosis.⁴⁵ A recent
273 study showed that that daily quercetin intake was associated with a lower prevalence
274 of T2DM in Chinese individuals, thus supporting a potential protective effect of
275 quercetin in the development of T2DM.⁴⁶ The possible anti-diabetic effects of
276 quercetin, which have been replicated both *in vivo* and *in vitro*, are linked mainly to
277 the anti-oxidant and anti-inflammatory actions of quercetin on pancreatic β cells.^{47,48}
278 Recent experimental data also suggested that quercetin might exert some beneficial
279 effects on risk of T2DM, possibly by inhibiting pancreatic β cell iron deposition and
280 ferroptosis.⁴⁵

281

282 **3.1.3 Other metabolic risk factors**

283 The newly proposed diagnostic criteria of MAFLD are based on the evidence of
284 hepatic steatosis (as assessed by histology, imaging techniques, or blood biomarkers),
285 combined with one of the following three conditions: overweight/obesity, T2DM, or
286 presence of metabolic dysregulation (i.e. defined by the presence of at least two of the
287 following seven metabolic risk abnormalities that are often also present with
288 metabolic syndrome: i.e. increased waist circumference, raised blood pressure, high
289 triglycerides, low HDL-cholesterol, increased plasma glucose concentration, IR or
290 elevated plasma C-reactive protein concentrations).^{4,49} Epidemiological studies have
291 also shown that elevated serum ferritin or iron overload are associated with higher
292 levels of plasma glucose, diastolic blood pressure, uric acid, and IR.^{50,51} Thus, as
293 schematically summarized in **Figure 1**, it is reasonable to hypothesize that ferroptosis
294 is closely related to obesity, T2DM or other coexisting metabolic disorders that occur
295 with MAFLD.

296

297 **3.2 Impact of ferroptosis on MAFLD severity**

298 **3.2.1 Hepatic inflammation or impaired liver function**

299 Iron overload may aggravate MAFLD/NAFLD by increasing the risk of hepatocyte
300 swelling, inflammation and fibrosis, thereby promoting the progression from simple
301 steatosis to steatohepatitis (NASH).^{52,53}

302

303 Recently, Tsurusaki et al. reported that ferroptosis is implicated in the development of
304 liver fat accumulation into NASH.⁵⁴ These authors also reported that inhibition of
305 ferroptosis prevents the development of NASH.⁵⁴ With development of NASH,
306 ferroptosis may lead to liver injury and inflammatory response, thereby providing a
307 new potential target for NASH treatment.⁵⁵

308

309 **3.2.2 Liver fibrosis**

310 Liver fibrosis is a complex pathophysiological process and an intermediate pathogenic
311 link in a variety of chronic liver diseases. Activation of HSC is a critical step in liver
312 fibrogenesis (**Figure 1**). Currently, liver transplantation is the only effective treatment
313 for patients with late-stage of liver fibrosis. However, the shortage of liver donors and
314 the risks of follow-up transplantation limit the treatment of patients with advanced
315 liver fibrosis. Therefore, it is urgent to find new treatment strategies for advanced
316 liver fibrosis.⁵⁶ The activation, proliferation, and transformation of HSCs are key
317 drivers in liver fibrogenesis. Therefore, focusing on HSCs as a target in MAFLD is
318 important in treatment strategies targeting liver fibrosis.

319

320 Iron is abundant in HSCs that is a pre-requisite condition for ferroptosis. From the
321 current evidence in animal studies (as discussed above), it appears that ferroptosis
322 might act as a two-edged mechanism in the development and progression of liver
323 fibrosis. Some studies showed that ferroptosis may exacerbate liver fibrosis and liver
324 injury.^{57,58} However, some experimental studies have recently suggested that
325 induction of ferroptosis could be also considered a new strategy to improve liver
326 fibrosis (as summarized in **Table 1**).⁵⁶⁻⁶² An animal study showed that artesunate
327 inhibits liver fibrogenesis through ferroptosis.⁵⁹ Magnesium isoglycyrrhizate may
328 regulate iron transport processes (mostly by up-regulating the expression of heme
329 oxygenase-1), and promote accumulation of Fe²⁺ and lipid peroxides RTMN induce
330 ferroptosis in HSC, thereby ameliorating liver fibrosis.⁶⁰ In addition, artemether up-
331 regulates the gene expression of P53, inhibits the SLC7A11 recombinant protein, and

332 inactivates GPX4 (eventually inducing ferroptosis of HSC), thereby inhibiting liver
333 fibrogenesis.⁶¹ Thus, ferroptosis might also inhibit the activation of HSC and thus
334 ameliorate liver fibrosis.

335

336 **Conclusions**

337 Ferroptosis is a newly discovered form of regulated cell death. Iron-dependent lipid
338 peroxidation is a major driver of ferroptosis and ferroptosis may also occur in
339 MAFLD. The concept of ferroptosis-inducing treatment regulating liver fibrosis is of
340 increasing interest in MAFLD. However, we suggest further mechanistic studies are
341 needed to better understand the role of ferroptosis in liver fibrogenesis. Experimental
342 evidence suggests that altered iron metabolism and lipid peroxidation are
343 pathophysiologically involved in the link between ferroptosis and MAFLD. However,
344 the role of ferroptosis in the pathophysiology of MAFLD is complex and needs
345 further investigation. On the one hand, ferroptosis can lead to the occurrence of
346 MAFLD and development of liver inflammation; whilst on the other hand, some
347 studies have shown that ferroptosis of HSC could inhibit the progression of liver
348 fibrosis. Further mechanistic studies of ferroptosis are required to better elucidate
349 whether altering this new form of regulated cell death has merit in the prevention and
350 treatment of MAFLD.

351

352 **References**

- 353 1. Yan HF, Zou T, Tuo QZ, et al. Ferroptosis: mechanisms and links
354 with diseases. *Signal transduction and targeted therapy*.
355 2021;6(1):49.
- 356 2. Stockwell BR, Friedmann Angeli JP, Bayir H, et al. Ferroptosis:
357 A Regulated Cell Death Nexus Linking Metabolism, Redox Biology,
358 and Disease. *Cell*. 2017;171(2):273–285.
- 359 3. Jiang X, Stockwell BR, Conrad M. Ferroptosis: mechanisms,
360 biology and role in disease. *Nat Rev Mol Cell Biol*.
361 2021;22(4):266–282.
- 362 4. Eslam M, Sanyal AJ, George J, International Consensus P. MAFLD:
363 A Consensus-Driven Proposed Nomenclature for Metabolic
364 Associated Fatty Liver Disease. *Gastroenterology*.
365 2020;158(7):1999–2014 e1991.

- 366 5. Eslam M, Newsome PN, Sarin SK, et al. A new definition for
367 metabolic dysfunction-associated fatty liver disease: An
368 international expert consensus statement. *J Hepatol.*
369 2020;73(1):202-209.
- 370 6. Zheng KI, Fan JG, Shi JP, et al. From NAFLD to MAFLD: a
371 "redefining" moment for fatty liver disease. *Chinese medical*
372 *journal.* 2020;133(19):2271-2273.
- 373 7. Mantovani A, Scorletti E, Mosca A, Alisi A, Byrne CD, Targher
374 G. Complications, morbidity and mortality of nonalcoholic fatty
375 liver disease. *Metabolism.* 2020;111S:154170.
- 376 8. Wang TY, George J, Zheng MH. Metabolic (dysfunction) associated
377 fatty liver disease: more evidence and a bright future.
378 *Hepatobiliary Surg Nutr* 2021.
- 379 9. Zheng KI, Eslam M, George J, Zheng MH. When a new definition
380 overhauls perceptions of MAFLD related cirrhosis care.
381 *Hepatobiliary Surg Nutr.* 2020;9(6):801-804.
- 382 10. Targher G, Tilg H, Byrne CD. Non-alcoholic fatty liver disease:
383 a multisystem disease requiring a multidisciplinary and
384 holistic approach. *The lancet Gastroenterology & hepatology.*
385 2021;6(7):578-588.
- 386 11. Dixon SJ, Lemberg KM, Lamprecht MR, et al. Ferroptosis: an
387 iron-dependent form of nonapoptotic cell death. *Cell.*
388 2012;149(5):1060-1072.
- 389 12. Zhang J, Li S, Li L, et al. Exosome and exosomal microRNA:
390 trafficking, sorting, and function. *Genomics Proteomics*
391 *Bioinformatics.* 2015;13(1):17-24.
- 392 13. Grignano E, Birsén R, Chapuis N, Bouscary D. From Iron
393 Chelation to Overload as a Therapeutic Strategy to Induce
394 Ferroptosis in Leukemic Cells. *Front Oncol.* 2020;10:586530.
- 395 14. Gautheron J, Gores GJ, Rodrigues CMP. Lytic cell death in
396 metabolic liver disease. *J Hepatol.* 2020;73(2):394-408.
- 397 15. He Z, Liao W, Song Q, et al. Role of ferroptosis induced by a
398 high concentration of calcium oxalate in the formation and
399 development of urolithiasis. *Int J Mol Med.* 2021;47(1):289-301.
- 400 16. Guo J, Xu B, Han Q, et al. Ferroptosis: A Novel Anti-tumor
401 Action for Cisplatin. *Cancer Res Treat.* 2018;50(2):445-460.
- 402 17. Zhao Y, Li Y, Zhang R, Wang F, Wang T, Jiao Y. The Role of
403 Erastin in Ferroptosis and Its Prospects in Cancer Therapy.
404 *OncoTargets and therapy.* 2020;13:5429-5441.
- 405 18. Yang WS, Stockwell BR. Ferroptosis: Death by Lipid
406 Peroxidation. *Trends Cell Biol.* 2016;26(3):165-176.
- 407 19. Bellanti F, Villani R, Facciorusso A, Vendemiale G, Serviddio
408 G. Lipid oxidation products in the pathogenesis of non-
409 alcoholic steatohepatitis. *Free Radic Biol Med.* 2017;111:173-

- 410 185.
- 411 20. Seki S, Kitada T, Yamada T, Sakaguchi H, Nakatani K, Wakasa K.
412 In situ detection of lipid peroxidation and oxidative DNA
413 damage in non-alcoholic fatty liver diseases. *J Hepatol*.
414 2002;37(1):56-62.
- 415 21. Serviddio G, Bellanti F, Tamborra R, et al. Uncoupling protein-
416 2 (UCP2) induces mitochondrial proton leak and increases
417 susceptibility of non-alcoholic steatohepatitis (NASH) liver to
418 ischaemia-reperfusion injury. *Gut*. 2008;57(7):957-965.
- 419 22. Koroglu E, Canbakan B, Atay K, et al. Role of oxidative stress
420 and insulin resistance in disease severity of non-alcoholic
421 fatty liver disease. *Turk J Gastroenterol*. 2016;27(4):361-366.
- 422 23. Zelber-Sagi S, Ivancovsky-Wajcman D, Fliss-Isakov N, et al.
423 Serum Malondialdehyde is Associated with Non-Alcoholic Fatty
424 Liver and Related Liver Damage Differentially in Men and Women.
425 *Antioxidants (Basel)*. 2020;9(7).
- 426 24. Britton LJ, Subramaniam VN, Crawford DH. Iron and non-alcoholic
427 fatty liver disease. *World journal of gastroenterology*.
428 2016;22(36):8112-8122.
- 429 25. Capelletti MM, Manceau H, Puy H, Peoc'h K. Ferroptosis in Liver
430 Diseases: An Overview. *Int J Mol Sci*. 2020;21(14).
- 431 26. Yang Z, Wu J, Li X, Xie D, Wang Y, Yang T. Association between
432 dietary iron intake and the prevalence of nonalcoholic fatty
433 liver disease: A cross-sectional study. *Medicine (Baltimore)*.
434 2019;98(43):e17613.
- 435 27. Pan X, Chen B, Liu W, et al. Circulating Iron Levels
436 Interaction with Central Obesity on the Risk of Nonalcoholic
437 Fatty Liver Disease: A Case-Control Study in Southeast China.
438 *Ann Nutr Metab*. 2019;74(3):207-214.
- 439 28. Rostoker G, Loridon C, Griuncelli M, et al. Liver Iron Load
440 Influences Hepatic Fat Fraction in End-Stage Renal Disease
441 Patients on Dialysis: A Proof of Concept Study. *EBioMedicine*.
442 2019;39:461-471.
- 443 29. Barrera C, Valenzuela R, Rincon MA, et al. Iron-induced
444 derangement in hepatic Delta-5 and Delta-6 desaturation
445 capacity and fatty acid profile leading to steatosis: Impact on
446 extrahepatic tissues and prevention by antioxidant-rich extra
447 virgin olive oil. *Prostaglandins Leukot Essent Fatty Acids*.
448 2020;153:102058.
- 449 30. Wang H, An P, Xie E, et al. Characterization of ferroptosis in
450 murine models of hemochromatosis. *Hepatology (Baltimore, Md)*.
451 2017;66(2):449-465.
- 452 31. Junge B, Carrion Y, Bosco C, et al. Effects of iron overload
453 and lindane intoxication in relation to oxidative stress,

- 454 Kupffer cell function, and liver injury in the rat. *Toxicol*
455 *Appl Pharmacol.* 2001;170(1):23-28.
- 456 32. Yamamoto M, Iwasa M, Iwata K, et al. Restriction of dietary
457 calories, fat and iron improves non-alcoholic fatty liver
458 disease. *Journal of gastroenterology and hepatology.*
459 2007;22(4):498-503.
- 460 33. Sakurai Y, Kubota N, Yamauchi T, Kadowaki T. Role of Insulin
461 Resistance in MAFLD. *Int J Mol Sci.* 2021;22(8).
- 462 34. Backe MB, Moen IW, Ellervik C, Hansen JB, Mandrup-Poulsen T.
463 Iron Regulation of Pancreatic Beta-Cell Functions and Oxidative
464 Stress. *Annu Rev Nutr.* 2016;36:241-273.
- 465 35. Gabrielsen JS, Gao Y, Simcox JA, et al. Adipocyte iron
466 regulates adiponectin and insulin sensitivity. *J Clin Invest.*
467 2012;122(10):3529-3540.
- 468 36. Chen X, Li J, Kang R, Klionsky DJ, Tang D. Ferroptosis:
469 machinery and regulation. *Autophagy.* 2020:1-28.
- 470 37. Ingold I, Berndt C, Schmitt S, et al. Selenium Utilization by
471 GPX4 Is Required to Prevent Hydroperoxide-Induced Ferroptosis.
472 *Cell.* 2018;172(3):409-422 e421.
- 473 38. Koruk M, Taysi S, Savas MC, Yilmaz O, Akcay F, Karakok M.
474 Oxidative stress and enzymatic antioxidant status in patients
475 with nonalcoholic steatohepatitis. *Ann Clin Lab Sci.*
476 2004;34(1):57-62.
- 477 39. Di Sessa A, Guarino S, Umamo GR, et al. MAFLD in Obese
478 Children: A Challenging Definition. *Children (Basel).*
479 2021;8(3).
- 480 40. Lee YS, Hwang LC, Hsu HY, Tsou MT. The Association Between
481 Different Obesity Phenotypes and Liver Fibrosis Scores in
482 Elderly Individuals with Fatty Liver in Taiwan. *Diabetes Metab*
483 *Syndr Obes.* 2021;14:1473-1483.
- 484 41. Yan HF, Liu ZY, Guan ZA, Guo C. Deferoxamine ameliorates
485 adipocyte dysfunction by modulating iron metabolism in ob/ob
486 mice. *Endocr Connect.* 2018;7(4):604-616.
- 487 42. Ma X, Pham VT, Mori H, MacDougald OA, Shah YM, Bodary PF. Iron
488 elevation and adipose tissue remodeling in the epididymal depot
489 of a mouse model of polygenic obesity. *PLoS one.*
490 2017;12(6):e0179889.
- 491 43. Fernández-Real JM, McClain D, Manco M. Mechanisms Linking
492 Glucose Homeostasis and Iron Metabolism Toward the Onset and
493 Progression of Type 2 Diabetes. *Diabetes Care.*
494 2015;38(11):2169-2176.
- 495 44. Heerspink HJL, Parving HH, Andress DL, et al. Atrasentan and
496 renal events in patients with type 2 diabetes and chronic
497 kidney disease (SONAR): a double-blind, randomised, placebo-

- 498 controlled trial. *Lancet (London, England)*.
499 2019;393(10184):1937–1947.
- 500 45. Li D, Jiang C, Mei G, et al. Quercetin Alleviates Ferroptosis
501 of Pancreatic β Cells in Type 2 Diabetes. *Nutrients*.
502 2020;12(10).
- 503 46. Yao Z, Gu Y, Zhang Q, et al. Estimated daily quercetin intake
504 and association with the prevalence of type 2 diabetes mellitus
505 in Chinese adults. *Eur J Nutr*. 2019;58(2):819–830.
- 506 47. Bardy G, Virsolvy A, Quignard JF, et al. Quercetin induces
507 insulin secretion by direct activation of L-type calcium
508 channels in pancreatic beta cells. *Br J Pharmacol*.
509 2013;169(5):1102–1113.
- 510 48. Eitah HE, Maklad YA, Abdelkader NF, Gamal El Din AA, Badawi MA,
511 Kenawy SA. Modulating impacts of quercetin/sitagliptin
512 combination on streptozotocin-induced diabetes mellitus in
513 rats. *Toxicol Appl Pharmacol*. 2019;365:30–40.
- 514 49. Zheng KI, Sun DQ, Jin Y, Zhu PW, Zheng MH. Clinical utility of
515 the MAFLD definition. *J Hepatol*. 2021;74(4):989–991.
- 516 50. Wu W, Yuan J, Shen Y, et al. Iron overload is related to
517 elevated blood glucose levels in obese children and aggravates
518 high glucose-induced endothelial cell dysfunction in vitro. *BMJ
519 Open Diabetes Res Care*. 2020;8(1).
- 520 51. Swaminathan S, Fonseca VA, Alam MG, Shah SV. The role of iron
521 in diabetes and its complications. *Diabetes Care*.
522 2007;30(7):1926–1933.
- 523 52. Sumida Y, Yoshikawa T, Okanou T. Role of hepatic iron in non-
524 alcoholic steatohepatitis. *Hepatol Res*. 2009;39(3):213–222.
- 525 53. Qi J, Kim JW, Zhou Z, Lim CW, Kim B. Ferroptosis Affects the
526 Progression of Nonalcoholic Steatohepatitis via the Modulation
527 of Lipid Peroxidation-Mediated Cell Death in Mice. *Am J Pathol*.
528 2020;190(1):68–81.
- 529 54. Tsurusaki S, Tsuchiya Y, Koumura T, et al. Hepatic ferroptosis
530 plays an important role as the trigger for initiating
531 inflammation in nonalcoholic steatohepatitis. *Cell Death Dis*.
532 2019;10(6):449.
- 533 55. Mao L, Zhao T, Song Y, et al. The emerging role of ferroptosis
534 in non-cancer liver diseases: hype or increasing hope? *Cell
535 Death Dis*. 2020;11(7):518.
- 536 56. Koyama Y, Xu J, Liu X, Brenner DA. New Developments on the
537 Treatment of Liver Fibrosis. *Dig Dis*. 2016;34(5):589–596.
- 538 57. Yu Y, Jiang L, Wang H, et al. Hepatic transferrin plays a role
539 in systemic iron homeostasis and liver ferroptosis. *Blood*.
540 2020;136(6):726–739.
- 541 58. Yang L, Wang H, Yang X, et al. Auranofin mitigates systemic

542 iron overload and induces ferroptosis via distinct mechanisms.
543 *Signal transduction and targeted therapy*. 2020;5(1):138.

544 59. Kong Z, Liu R, Cheng Y. Artesunate alleviates liver fibrosis by
545 regulating ferroptosis signaling pathway. *Biomed Pharmacother*.
546 2019;109:2043-2053.

547 60. Sui M, Jiang X, Chen J, Yang H, Zhu Y. Magnesium
548 isoglycyrrhizinate ameliorates liver fibrosis and hepatic
549 stellate cell activation by regulating ferroptosis signaling
550 pathway. *Biomed Pharmacother*. 2018;106:125-133.

551 61. Wang L, Zhang Z, Li M, et al. P53-dependent induction of
552 ferroptosis is required for artemether to alleviate carbon
553 tetrachloride-induced liver fibrosis and hepatic stellate cell
554 activation. *IUBMB Life*. 2019;71(1):45-56.

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577 **Table Legend**

578 **Table 1.** Experimental studies examining the role of ferroptosis in liver fibrosis.

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580 **Figure Legend**

581 **Figure 1.** Potential regulatory mechanisms of ferroptosis in MAFLD.

582 Ferroptosis is closely related to obesity, type 2 diabetes mellitus (T2DM) and other
583 metabolic risk factors, all of which occur in MAFLD. The main regulatory
584 mechanisms of ferroptosis in MAFLD include increased lipid peroxidation and iron
585 overload. Induction of ferroptosis of hepatocytes may also inhibit fibrogenesis and
586 serve as potential anti-fibrotic treatment. Moreover, ferritinophagy-mediated hepatic
587 stellate cells (HSC)-ferroptosis may be also responsible for its anti-fibrotic efficacy.

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