Early Change in the Plasma Levels of Circulating Soluble Immune Checkpoint Proteins in Patients with Unresectable Hepatocellular Carcinoma Treated by Lenvatinib or Transcatheter Arterial Chemoembolization

Naoshi Odagiri, Hoang Hai, Le Thi Thanh Thuy, Minh Phuong Dong, Maito Suoh, Kohei Kotani, Atsushi Hagihara, Sawako Uchida-Kobayashi, Akihiro Tamori, Masaru Enomoto, Norifumi Kawada

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Highlights
◇肝がん患者において、既存治療である血管新生阻害剤またはTACEなどが、早期にがん微小環境の免疫状態に影響を与えていることを明らかにした。
◇肝がんでは、免疫チェックポイント阻害剤に既存治療を併用することで、相乗効果が得られることが期待される。

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| 概要 |
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| レンバチニブという血管新生阻害剤または肝動脈化学栓塞術（TACE）で治療を受けた肝がん患者の血液で、16種類の可溶性免疫チェックポイント分子を測定したところ、治療導入の1週間後に早くも複数の分子の濃度が有意に変化していることを発見しました。この発見は、肝がんに対しては既存の治療法であっても、がん微小環境の免疫状態に影響を与えている可能性を示しています。
肝がんは、日本で年間に3万人近くが発生する重要な疾患であり、治療法について研究が進められています。がん免疫療法である免疫チェックポイント阻害剤は様々ながんで使われ効果をあげていますが、肝がんにおいては単剤での効果は認められていません。今回の研究成果より、肝がんに対しては免疫チェックポイント阻害剤に血管新生阻害剤またはTACEなどの既存治療を併用することで相乗効果が得られる可能性が期待されます。 |

| Description |
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| ＜研究の背景＞
従来から肝がんに対する治療法として、外科手術、ラジオ波焼灼療法、肝動脈化学栓塞術（TACE）などが行われてきました。これに加え全身化学療法として、2009年にソラフェニブが承認されて以来、レゴラフェニブ、レンバチニブ、ラムシルマブなど、がん細胞が増殖するために必要な血流を遮断する血管新生阻害剤が使われています。
一方、近年はがん免疫療法も目覚ましく進歩しています。一例として、2018年にノーベル薬学賞を授与された京都大学の本庶佑名誉教授と米テキサス大学のジェームズ・アリソン教授は、免疫の司令塔であるTリンパ球表面にPD-1またはCTLA-4という分子を発見し、これをもとにオプジーボ®など免疫チェックポイント阻害剤が開発されました。
残念ながら肝がんに対して免疫チェックポイント阻害剤は承認されていませんが、血管新生阻害剤やTACEなど既存治療を併用することで相乗効果が得られ、今後、第一選択薬になる可能性があります。
こうした既存治療と免疫チェックポイント阻害剤の併用治療については臨床試験でも有望な結果が出ている一方、これら既存治療ががん微小環境の免疫状態にどのように影響しているのかについては不明な部分が多いのが現状です。

＜研究の内容＞
研究チームは、血管新生阻害剤レンバチニブまたはTACEで治療を受けた肝がん患者さんの血液で、PD-1、CTLA-4をはじめ16種類の可溶性免疫チェックポイント分子の濃度を測定しました。
その結果、①レンバチニブで治療を行った患者さんでは、1週間後に早くもsCD27が有意に低下、sCD40、sTIM-3が有意に上昇していることが分かりました（図1）。また、各免疫チェックポイント分子の変化に関する相関を解析したところ、sCTLA-4とsCD86/sCD80、sPD-1とsPD-L1など、受容体とリガンドの関係にある分子同士で相関がみられました（図2）。
これらの結果は、血管新生阻害剤であるレンバチニブが、がん微小環境の免疫状態に影響している可能性を示しています。②TACE治療例でも、sCD27、sCD40、sTIM-3含む8種類の可溶性免疫チェックポイント分子の濃度が変化していることも分かりました（図
この結果は、TACE もまたがん微小環境の免疫状態に影響を与える可能性を示しています。

図 1. レンバチニブ治療開始 1 週後の肝がん患者さんの血液中の免疫チェックポイント分子の変化
治療開始早期の段階で sCD27 は有意に低下し、sCD40、sTIM-3 は有意に上昇していることが分かりました。

図 2. レンバチニブ治療開始 1 週後の肝がん患者さんの血液中の免疫チェックポイント分子間の変化の相関
sCTLA-4 と sCD-80/sCD-86、sPD-1 と s PD-L1 など、受容体とリガンドの関係にある分子同士の変化に相関がみられました。

図 3. TACE 治療 1 週後の肝がん患者さんの血液中の免疫チェックポイント分子の変化
上図の 4 分子を含む 8 種類の可溶性免疫チェックポイント分子の濃度が有意に変化していることが分かりました。

＜今後の展開＞
研究チームでは以前に血管新生阻害剤であるソラフェニブでも同様の解析を行い、ソラフェニブががん微小環境の免疫状態に影響を与えることを示しています。今回の結果からレンバチニブや TACE による治療もがん微小環境への作用を有していることが分かりました。これらの治療法ががん微小環境修飾機序をより詳細に解析することで、今後、血管新生阻害剤や TACE と免疫チェックポイント阻害剤を併用した、より効果的な肝がん治療法が開発できる可能性が考えられます。
免疫チェックポイント阻害剤は一部の患者さんに著効が得られる反面、その効果が予測
しにくいこと、重篤な副作用がみられる場合があること、薬価が高いことなどが問題になっています。先行研究では、特定の分子の変動が患者さんの予後予測に有用であったことも見出しています※1。研究チームでは今回の研究成果を足がかりに、より効果的な血管新生阻害剤やTACEと免疫チェックポイント阻害剤との併用治療の開発、より精度が高く治療効果を予測できるマーカーの開発につなげたいとしています。

※1 Dong MP, Enomoto M, Thuy LTT, et al. Clinical significance of circulating soluble immune checkpoint proteins in sorafenib-treated patients with advanced hepatocellular carcinoma. Sci Rep. 2020;10(1):3392. Published 2020 Feb 25. doi:10.1038/s41598-020-60440-5

‘肝がん既存治療のメカニズムの一端を解明：将来の免疫チェックポイント阻害剤との併用治療に期待！’ 大阪市立大学. https://www.osaka-cu.ac.jp/ja/news/2020/200805. （参照 2020-08-05）
Early Change in the Plasma Levels of Circulating Soluble Immune Checkpoint Proteins in Patients with Unresectable Hepatocellular Carcinoma Treated by Lenvatinib or Transcatheter Arterial Chemoembolization

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Abstract: Immune checkpoint inhibitors, combined with anti-angiogenic agents or locoregional treatments (e.g., transarterial chemoembolization (TACE)), are expected to become standard-of-care for unresectable hepatocellular carcinoma (HCC). We measured the plasma levels of 16 soluble checkpoint proteins using multiplexed fluorescent bead-based immunoassays in patients with HCC who underwent lenvatinib (n = 24) or TACE (n = 22) treatment. In lenvatinib-treated patients, plasma levels of sCD27 (soluble cluster of differentiation 27) decreased (p = 0.040) and levels of sCD40 (p = 0.014) and sTIM-3 (p < 0.001) were increased at Week 1, while levels of sCD27 (p < 0.001) were increased significantly at Weeks 2 through 4. At Week 1 of TACE, in addition to sCD27 (p = 0.028), sCD40 (p < 0.001), and sTIM-3 (soluble T-cell immunoglobulin and mucin domain–3) (p < 0.001), levels of sHVEM (soluble herpesvirus entry mediator) (p = 0.003), sTLR-2 (soluble Toll-like receptor 2) (p = 0.009), sCD80 (p = 0.036), sCTLA-4 (soluble cytotoxic T-lymphocyte antigen 4) (p = 0.005), sGITR (soluble glucocorticoid-induced tumor necrosis factor receptor) (p = 0.030), sGITRL (soluble glucocorticoid-induced TNFR-related ligand) (p = 0.090), and sPD-L1 (soluble programmed death-ligand 1) (p = 0.070) also increased. The fold-changes in soluble checkpoint receptors and their ligands, including sCTLA-4 with sCD80/sCD86 and sPD-1 (soluble programmed cell death domain–1) with sPD-L1 were positively correlated in both the lenvatinib and TACE treatment groups. Our results suggest that there are some limited differences in immunomodulatory effects between anti-angiogenic agents and TACE. Further studies from multicenters may help to identify an effective combination therapy.

Keywords: HCC; liver cancer; molecular-targeted agent; TACE; tyrosine kinase inhibitor

1. Introduction

Primary liver cancer, of which hepatocellular carcinoma (HCC) is the most common type, is the third most common cause of cancer-related deaths worldwide [1,2]. Patients with early-stage HCC can potentially receive curative treatments, such as surgical resection, transplantation, or ablation, and patients at the intermediate stage can undergo transarterial chemoembolization (TACE), but those at an advanced stage of disease are only likely to benefit from systemic therapies [3–5]. Molecular-targeted
therapies, the two anti-angiogenic tyrosine kinase inhibitors sorafenib [6] and lenvatinib [7] are used as first-line treatments for patients with advanced-stage HCC. Meanwhile, other anti-angiogenic agents like regorafenib [8], cabozantinib [9], and ramucirumab (if α-fetoprotein > 400 ng/mL) [10] have been licensed as second-line treatments. Immune checkpoint inhibitors, such as anti–PD-1 (programmed cell death domain–1) pembrolizumab [11], nivolumab [12] or nivolumab combined with anti–CTLA-4 (cytotoxic T-lymphocyte antigen 4) (ipilimumab) therapy [13], may also be indicated for sorafenib-refractory patients [14–16].

However, therapeutic responses to immunotherapy alone are obtained in a minority of patients with HCC. Current clinical trials are therefore focusing on whether combinations of different types of treatments, including anti-angiogenic agents and TACE, may be promising for the enhancement of the antitumor effects of immune checkpoint inhibitors [17]. For example, the combinations of lenvatinib plus pembrolizumab [18] and bevacizumab plus anti–PD-L1 atezolizumab therapy [19] are currently being tested in clinical investigations. In addition, tremelimumab, an anti–CTLA-4 therapy, was more effective when paired with TACE than as monotherapy [20].

In our previous study [21], sorafenib treatment in patients with unresectable HCC provoked dynamic changes in soluble immune checkpoint protein levels as revealed using multiplexed fluorescent bead-based immunoassays. To date, circulating soluble checkpoint proteins, which are part of a family of full-length receptors produced by messenger RNA expression or by the cleavage of membrane-bound proteins, have been studied extensively in various cancers, but not in HCC [22]. Further, changes in the plasma levels of soluble proteins during the early days of treatment with other anti-angiogenic agents or locoregional treatments (e.g., TACE) have yet to be determined.

We hypothesized that lenvatinib and TACE therapies as well as sorafenib would affect plasma levels of soluble immune checkpoint proteins. An understanding of the effects of these therapies on soluble immune checkpoint proteins may provide insight into their immunomodulative characteristics, which would help to develop more effective combination immunotherapies for HCC. In this study, we measured the concentrations of 16 soluble immune checkpoint proteins (Table A1) in plasma samples obtained over four weeks of treatment from patients with unresectable HCC who underwent lenvatinib or TACE treatment.

2. Results

2.1. Patient Characteristics

The baseline characteristics of patients with HCC in the lenvatinib (n = 24) and TACE (n = 22) groups are described in Table 1. In brief, the median age of patients was about 75 years (75 vs. 76 years; p = 0.530) and males accounted for the majority (75.0% vs. 68.2%; p = 0.746) of the population in both treatment groups, with no significant difference apparent between the two groups. Most patients in the TACE group had Barcelona Clinic Liver Cancer (BCLC) [23] stage A (54.5%) cancer, while all patients had BCLC stage B (54.2%) or stage C (45.8%) cancer in the lenvatinib group (p < 0.001). Some patients in the TACE group underwent the therapy as their initial HCC treatment, but none in the lenvatinib group (36.4% vs. 0%; p = 0.001). No significant differences were found between the lenvatinib and TACE groups with respect to the etiology of liver disease, Eastern Cooperative Oncology Group performance status score, aspartate and alanine aminotransferase (AST and ALT) levels, gamma-glutamyl transferase level, hepatic reserve (Child–Pugh class or albumin–bilirubin score [24,25]) and α-fetoprotein and des-γ-carboxy prothrombin levels as tumor markers.
Table 1. Characteristics of hepatocellular carcinoma (HCC) patients at baseline.

| Characteristics          | Lenvatinib (n = 24) | TACE (n = 22) | p Value |
|--------------------------|---------------------|---------------|---------|
| Age                      | 75 (69, 78)         | 76 (69, 80)   | 0.530   |
| Sex                      | Male 18 (75.0)      | 15 (68.2)     | 0.746   |
|                         | Female 6 (25.0)     | 7 (31.8)      |         |
| Etiology                 | Alcohol 5 (20.8)    | 3 (13.6)      | 0.536   |
|                         | HBV 3 (12.5)        | 1 (4.5)       |         |
|                         | HCV 10 (41.7)       | 13 (59.1)     |         |
|                         | HBV + HCV 1 (4.2)   | 0 (0.0)       |         |
|                         | NASH 1 (4.2)        | 3 (13.6)      |         |
|                         | Unknown 4 (16.7)    | 2 (9.1)       |         |
| ECOG Performance Status  | 0 or 1 23 (95.8)    | 19 (86.4)     | 0.336   |
|                         | 2 1 (4.2)           | 3 (13.6)      |         |
| Aspartate aminotransferase| 39 (27, 59)        | 39 (30, 50)   | 0.826   |
| Alanine aminotransferase | 27 (20, 52)         | 23 (16, 46)   | 0.567   |
| Gamma-glutamyl transferase| 62 (30, 122)       | 61 (32, 120)  | 0.930   |
| Child-Pugh class         | A 21 (87.5)         | 18 (81.8)     | 0.694   |
|                         | B 3 (12.5)          | 4 (18.2)      |         |
| ALBI grade               | 1 8 (33.3)          | 8 (36.4)      | 0.999   |
|                         | 2 16 (66.7)         | 14 (63.6)     |         |
| α-Fetoprotein            | 59.4 (11.4, 1123.7) | 21.2 (5.3, 55.9) | 0.126 |
| Des-γ-carboxy prothrombin| 105 (72, 1312)    | 137 (56, 282) | 0.605   |
| BCLC stage               | A 0 (0.0)           | 12 (54.5)     | <0.001  |
|                         | B 13 (54.2)         | 7 (31.8)      | 0.418   |
|                         | C 11 (45.8)         | 3 (13.6)      |         |
| Previous therapies       | None 0 (0.0)        | 8 (36.4)      | 0.001   |
|                         | Resection 5 (20.8)  | 2 (9.1)       | 0.418   |
|                         | RFA/PEI 11 (45.8)   | 7 (31.8)      | 0.378   |
|                         | TACE 23 (95.8)      | 9 (40.9)      | <0.001  |
|                         | HAIC 3 (12.5)       | 3 (13.6)      | 0.999   |
|                         | Radiation 2 (8.3)   | 0 (0.0)       | 0.490   |
|                         | Chemotherapy 0 (0.0)| 2 (9.1)       | 0.223   |

Data are shown as median [interquartile range] or number (%). Abbreviations: HBV, hepatitis B virus; HCV, hepatitis C virus; NASH, non-alcoholic steatohepatitis; ECOG, Eastern Cooperative Oncology Group; ALBI, albumin–bilirubin; BCLC, Barcelona Clinic Liver Cancer; RFA, radiofrequency ablation; PEI, percutaneous ethanol injection; TACE, transarterial chemoembolization; HAIC, hepatic arterial infusion chemotherapy.

2.2. Changes in Plasma Soluble Checkpoint Protein Levels at Week 1 after the Initiation of Lenvatinib

First, we investigated whether lenvatinib affects the plasma levels of immune checkpoint proteins in the early phase of treatment. As previously reported [21], we analyzed soluble checkpoint protein levels in patients with HCC at Week 1 of lenvatinib treatment. Ultimately, a significant decrease was observed in the level of soluble cluster of differentiation 27 (sCD27) (p = 0.040) and significant increases were found in the levels of sCD40 (p = 0.014) and soluble T-cell immunoglobulin and mucin domain–3 (sTIM-3) (p < 0.001) when compared with at baseline. Meanwhile, no significant changes were found in soluble herpesvirus entry mediator (sHVEM) and the other 12 immune checkpoint proteins (Figures 1 and A1).
while the other 12 immune checkpoint proteins showed similar outcomes (Appendix A Figure A2).

In addition to sCD27 (soluble CD27), we also investigated patients who underwent PEI, percutaneous ethanol injection; TACE, transarterial chemoembolization; HAIC, hepatic arterial chemoembolization. We analyzed soluble checkpoint protein levels in the plasma of patients with HCC at 1 week after TACE. In addition to sCD27, soluble T-cell immunoglobulin and mucin domain–3 (sTIM-3), and glucocorticoid-induced TNFR-related ligand (sGITRL) (p = 0.070 and 0.090). Also, the change in sTIM-3 was no longer significant at this time point (Figure 2), while the other 12 immune checkpoint proteins showed similar outcomes (Appendix A Figure A2).

Figure 1. Changes in soluble immune checkpoint protein levels in plasma after 1 week of lenvatinib treatment. Box plots show the sCD27, sCD40, sTIM-3, and sHVEM levels in HCC patients at baseline and Week 1 of treatment. The vertical lengths of the boxes indicate the interquartile ranges and the lines in the boxes suggest the median values, while the error bars show the minimum and maximum values (in a range). The Wilcoxon signed-rank test was used. HCC, hepatocellular carcinoma; sTIM-3, soluble T-cell immunoglobulin and mucin domain–3; sHVEM, soluble herpesvirus entry mediator; sCD, soluble cluster of differentiation.

2.3. Changes in Plasma Soluble Checkpoint Protein Levels at Weeks 2 through 4 after the Initiation of Lenvatinib

Next, we sought to reveal changes in immune checkpoint protein levels at the later stage of lenvatinib therapy. We analyzed soluble checkpoint protein levels in the plasma of patients with HCC at Weeks 2 through 4 of lenvatinib treatment. The increase in sCD27 level was significant (p < 0.001). A trend toward increasing sCD40 and sHVEM levels was also observed, but no change reached statistical significance (p = 0.070 and 0.090). Also, the change in sTIM-3 was no longer significant at this time point (Figure 2), while the other 12 immune checkpoint proteins showed similar outcomes (Appendix A Figure A2).

Figure 2. Changes in soluble immune checkpoint protein levels in plasma after 2 to 4 weeks of lenvatinib treatment. Box plots demonstrate the sCD27, sCD40, sTIM-3, and sHVEM levels in HCC patients at baseline and Weeks 2 through 4 of treatment. The vertical lengths of the boxes indicate the interquartile ranges and the lines in the boxes display the median values, while the error bars show the minimum and maximum values (in a range). The Wilcoxon signed-rank test was used.

2.4. Changes in Plasma Soluble Checkpoint Protein Levels at Week 1 after TACE

Lenvatinib can suppress tumor blood flow by its pharmacological anti-angiogenic effects. To establish a contrast with those who received lenvatinib, we also investigated patients who underwent conventional TACE, which disrupts tumor blood flow via the artificial embolization of hepatic arteries. We analyzed soluble immune checkpoint protein levels in the plasma of patients with HCC at 1 week after TACE. In addition to sCD27 (p = 0.028), sCD40 (p < 0.001), sTIM-3 (p < 0.001), and sHVEM (p = 0.003) levels, which exhibited significant (or marginal) changes at Week 1 or Weeks 2 through 4 of lenvatinib treatment, the levels of another six proteins—namely, soluble Toll-like receptor 2 (sTLR-2) (p = 0.009), sCD80 (p = 0.036), sCTLA-4 (soluble cytotoxic T-lymphocyte antigen 4) (p = 0.005), soluble glucocorticoid-induced TNFR-related protein (sGITR) (p = 0.030), soluble glucocorticoid-induced TNFR-related ligand (sGITRL) (p = 0.090), and sPD-L1 (soluble programmed death-ligand 1) (p = 0.070)—were also increased (Figure 3). However, the levels of the remaining six immune checkpoint proteins showed no significant changes (Appendix A Figure A3).
2.5. Relationships between Fold-Changes in Plasma Soluble Immune Checkpoint Protein Levels

The correlations between the fold-changes in the soluble forms of immune checkpoint proteins in plasma at Week 1 of lenvatinib treatment are shown in Figure 4. The fold-changes in soluble checkpoint receptors and their ligands, including sCTLA-4 with sCD80 (p < 0.001; r = 0.82)/sCD86 (p < 0.001; r = 0.78), and sPD-1 with sPD-L1 (p < 0.001; r = 0.91), were positively correlated (Figure 4a). Among the three soluble checkpoint proteins with significant change at Week 1, sCD40 was positively correlated with some proteins in fold-changes; sCD86 (p = 0.005; r = 0.62), sPD-1 (p = 0.010; r = 0.58), sPD-L1 (p = 0.046; r = 0.48) (Figure 4b), sCD28 (p = 0.012; r = 0.57), sTLR-2 (p = 0.048; r = 0.46), and sHVEM (p = 0.042; r = 0.47).

At Weeks 2 to 4 of lenvatinib treatment, the fold-changes in soluble immune checkpoint receptors and their ligands in plasma, including sCTLA-4 with sCD80 (p = 0.055; r = 0.55)/sCD86 (p = 0.009; r = 0.71) and sPD-1 with sPD-L1 (p < 0.001; r = 0.86), were also positively correlated (Appendix A Figure A4a). Separately, the fold-change in sCD27 was positively correlated with those of sCD86 (p = 0.030; r = 0.61), sPD-1 (p = 0.010; r = 0.70), and sPD-L1 (p = 0.050; r = 0.56), respectively (Appendix A Figure A4b).

The correlations between the fold-changes in soluble forms of immune checkpoint proteins in plasma at Week 1 of TACE are shown in Figure 5. Again, the fold-changes in soluble checkpoint receptors and their ligands, including sCTLA-4 with sCD80 (p = 0.018; r = 0.50)/sCD86 (p < 0.001; r = 0.72) and sPD-1 with sPD-L1 (p < 0.001; r = 0.89), were positively correlated (Figure 5a). A strong correlation was noted between the fold-changes in the sCD80 and sTLR-2 levels (p < 0.001; r = 0.89), sCD40 and sHVEM levels (p < 0.001; r = 0.78), and sHVEM and sTIM-3 levels (p < 0.001; r = 0.72), respectively (Figure 5b).
Figure 4. The relationships between fold-changes in soluble immune checkpoint protein levels in plasma at 1 week of lenvatinib treatment are presented. (a) A positive correlation was found between fold-changes in soluble checkpoint receptors and their ligands, including sCTLA-4 with sCD80/sCD86 and sPD-1 with sPD-L1, respectively. (b) sCD40 was positively correlated with some proteins in fold-changes, including sCD86, sPD-1, and sPD-L1. Spearman’s rank correlation test was used. A p-value of less than 0.05 was considered to be statistically significant.

Figure 5. The relationships between fold-changes in soluble immune checkpoint protein levels in plasma at 1 week of receiving TACE are shown. (a) A positive correlation was found between the fold-changes of soluble checkpoint receptors and their ligands, including sCTLA-4 with sCD80/sCD86 and sPD-1 with sPD-L1. (b) Separately, a strong correlation was observed between the fold-changes in sCD80 and sTLR-2 levels, sCD40 and sHVEM levels, and sHVEM and sTIM-3 levels. Spearman’s rank correlation test was used. A p-value of less than 0.05 was considered to be statistically significant.
3. Discussion

This study attempted the simultaneous quantification of 16 soluble immune checkpoint proteins in the plasma of patients with HCC who were observed during the early phase of treatment with lenvatinib or TACE. These immune checkpoint proteins include soluble forms of stimulatory or inhibitory factors, which modulate T-cell activation/proliferation and compose the cancer-immunity cycle [26]. In our previous report [21], we examined changes to these immune checkpoint proteins in sorafenib-treated HCC. The current results could offer additional data to compare changes in soluble checkpoint protein levels among tyrosine kinase inhibitors and TACE, which may clarify the immunomodulatory aspects of these treatments.

In this study, the plasma level of sCD27 was decreased and those of sCD40 and sTIM-3 were increased significantly at Week 1, while the level of sCD27 was increased significantly at Weeks 2 through 4 of lenvatinib treatment (Figures 1 and 2). Although CD27 and CD40 are stimulatory factors and TIM-3 is an inhibitory factor, respectively, in the cancer-immunity cycle, soluble forms of stimulatory/inhibitory factors do not necessarily have set positive/negative immune effects; currently, the functions of these soluble proteins are yet to be fully defined. In short, (1) CD27 supports the antigen-specific expansion of naïve T-cells and is vital for the generation of T-cell memory [27]. A previous study found that circulating sCD27 constitutes a functional protein directly involved in CD8$^+$ T-cell activation [28]. A persistent increase in the sCD27 level during Weeks 2 through 4 of lenvatinib treatment may reflect the CD8$^+$ T-cell–related immunomodulatory effect of lenvatinib, which is in line with a previous study [29] revealing that the antitumor activity of lenvatinib was significantly diminished by CD8$^+$ T-cell depletion. Further, (2) CD40 plays an important role mainly in the signaling pathways for the functioning of B-cells, monocytes, and dendritic cells [30]. In recent research [31], plasma sCD40 levels were upregulated in correlation with disease severity in patients with alcoholic hepatitis, indicating the existence of dysregulation of the immune system in chronic liver disease. Also, (3) TIM-3 promotes the exhaustion of T-cells in various types of cancer [32]. One study suggests circulating sTIM-3 might competitively bind to galectin-9 (a ligand of TIM-3), preventing a TIM-3/galectin-9–mediated immune response [33]. The increase in plasma sTIM-3 during lenvatinib treatment may restore immune exhaustion.

In vitro and in vivo preclinical studies have provoked the thoughts that there may be different immunomodulatory effects among similar tyrosine kinase inhibitors [34]. Lenvatinib may decrease tumor-associated macrophages, facilitate polarization from an M2-like phenotype toward an M1-like phenotype, and enhance CD4$^+$ and CD8$^+$ T-cell tumor infiltration, while sorafenib may have the opposite effects [35]. In our previous study of sorafenib [21], 11 of 16 soluble immune checkpoint proteins—most of which were inhibitory factors—experienced significant increases in plasma at two weeks of treatment. In this study, sCD27 displayed a significant change at two weeks. Both sorafenib and lenvatinib basically inhibit vascular endothelial growth factor receptors (VEGFRs) 1 through 3, fibroblast growth factor receptors (FGFRs) 1 through 4, platelet-derived growth factor receptor (PDGFR)-α, RET, and KIT. One potential explanation for the discrepancy is the existence of variable inhibitory profiles among the drugs against VEGFRs and FGFRs; specifically, lenvatinib is known to show more potent inhibitory activities than sorafenib against these receptor tyrosine kinases [36]. VEGFRs are particularly important because they activate various key components such as regulatory T-cells, tumor-associated macrophages, and myeloid-derived suppressor cells. Cytokines released by these cells inhibit natural killer cell activation and CD8$^+$ T-cell proliferation, driving the emergence of an immunosuppressive microenvironment [37].

The tumor microenvironment actively participates in drug-resistance acquisition [38] in both primary and metastatic lesions [39] of HCC and other solid tumors [40]. Previous studies reported that an inflammatory microenvironment, circulating immune cells, and cytokines etc. play a significant role in the prognosis of HCC [41,42]. For example, the B-type Raf kinase (BRAF) mutation, one of the prognostic factors, could play a role in the response to tyrosine kinase inhibitors [43]. However, we found no significant difference in changes in soluble immune checkpoint protein levels between groups classified according to the response to sorafenib [21] or lenvatinib treatment, possibly due
to the small sample size. Further investigation is required in a large group of patients to establish a determinant in driving clinical decision-making, which today are an unmet clinical need and a challenge for immunotherapy.

As part of our research, we also analyzed the levels of soluble immune checkpoint proteins in TACE-treated patients. Hepatic arterial embolization induces tumor necrosis and focal inflammation by blocking the tumor blood supply, which can impact cancer immunity by creating a source of tumor-associated antigens and enhancing the tumor-specific T-cell response [44]. Tampaki et al. [45] reported that TACE provokes a significant increase in the sTIM-3 level in plasma within the first week posttreatment, suggesting a reactive expansion of TIM-3 expression by T-cells as a negative feedback mechanism in response to intense immune stimulation following tumor necrosis. In our research encompassing a comprehensive measurement of 16 immune checkpoints, not only did the level of sTIM-3 but also those of many more soluble proteins changed significantly in the first week after TACE (Figure 3). However, some confounding factors may accompany TACE. When compared with systemic therapies, TACE presents a greater likelihood of inducing more sudden hypoxia in treated lesions and produces more hypoxia-related factors, which can influence components of cancer-immunity in the short-term. Therefore, the interpretation of results among TACE-treated patients is difficult and further research is necessary to better understand the treatment’s immunomodulatory effect.

Correlation analyses involving two soluble immune checkpoint proteins in plasma revealed that the fold-changes in soluble receptors and their ligands were positively correlated in both lenvatinib-treated and TACE-treated patients (Figures 4a and 5a), suggesting that multiplexed fluorescent immunoassays are capable of accurate and simultaneous quantification of small amounts of the proteins. More interestingly, sCD40 at 1 week and sCD27 at two to four weeks, respectively, were positively correlated with sCTLA-4, sCD86, sPD-1, and sPD-L1 (Figure 4b and Appendix A Figure A4b). While the mechanisms regulating these proteins remain unknown, sCD40 and sCD27 may be upregulated in cooperation with major checkpoint molecules such as sCTLA-4, sCD86, sPD-1, and sPD-L1. At Week 1 of TACE treatment, many more checkpoint pairs showed correlations unexpectedly (Figure 5b). For example, CD80 and TLR-2 have both been generally used as specific M1 macrophage surface markers in previous research [46]. Elsewhere, CD40 and HVEM messenger RNAs were coexpressed in the bioinformatic analysis of bladder cancer in the Cancer Genome Atlas database [47]. HVEM is a ligand of B- and T-lymphocyte attenuator (BTLA) and soluble BTLA in combination with sTIM-3 may be able to predict the rates of disease recurrence and survival among renal cell cancer patients [48].

Our study has several limitations. First, this is not a prospective study but instead a retrospective observational study. Moreover, randomization was not performed. In current guidelines [3–5], TACE is essentially indicated as a treatment option for patients with intermediate-stage HCC, while systemic therapy is suggested for those with advanced-stage HCC. Along these lines, in this study, the baseline characteristics of patients in the lenvatinib and TACE groups were not similar to one another (Table 1). It is therefore difficult to accurately compare the changes in soluble immune checkpoint levels in plasma between these treatment groups. Second, we could not collect plasma samples at Week 2 through 4 after TACE because of the less frequent hospital visits made by TACE-treated patients. We observed dynamic changes in soluble checkpoint levels in plasma at Week 1 of TACE but could not determine how long they persisted. Third, the sample size was small. Sample size affects the power or ability of all statistical tests to detect a relationship between two variables when it truly exists. The limitations of statistical analyses, such as the Wilcoxon signed-rank test and Spearman’s rank correlation test, are equally worth noting [49]. Lastly, we could not investigate associations between changes in soluble checkpoints and the outcomes of HCC patients because a majority of the included patients are still alive. Further studies are needed to clarify the clinical significance of circulating soluble checkpoint proteins in patients with unresectable HCC.
4. Materials and Methods

4.1. Patients

Between May 2018 and February 2020, we initiated lenvatinib treatment in 55 patients (46 males and nine females; median age: 73 (range: 42–85) years) with unresectable HCC at our institute. In this retrospective cohort study, 24 patients who received lenvatinib as a first-line systemic therapy (18 males and six females; median age: 75 (range: 55–89) years) were included. Lenvatinib therapy was principally indicated for patients with a good performance status and compensated liver disease for whom locoregional therapies were not indicated either due to the presence of vascular invasion, distant metastasis, or a TACE-refractory state, in accordance with available guidelines [3–5]. Lenvatinib (Lenvima®; Eisai Co. Ltd., Tokyo, Japan) was orally given to patients with unresectable HCC. The dose of lenvatinib was determined based on body weight, with initial dosages of 12 mg/day and 8 mg/day given to those over 60 kg and those under 60 kg, respectively. Plasma samples were collected at baseline ($n = 24$), during the first week (days 3–7 after initiating therapy; $n = 19$), and during the second to fourth weeks (days 8–28 after initiating therapy; $n = 13$).

As part of an effort to compare with lenvatinib treatment, we also studied 22 patients (15 males and seven females; median age: 76 (range: 44–86) years) who underwent conventional TACE during the same period. TACE was principally indicated for those with three or less HCC masses measuring greater than 3 cm or for four or more HCC masses in patients with Child–Pugh A/B hepatic reserve without extrahepatic metastasis or vascular invasion, in accordance with available guidelines [3–5]. Conventional TACE consisted of intra-arterial injection of lipiodol plus epirubicin followed by the injection of an embolic agent, Gelpart (Nippon Kayaku Co. Ltd., Tokyo, Japan), to interrupt blood flow. Plasma samples were collected at baseline and during the first week (days 3–7 after initiating therapy).

4.2. Soluble Immune Checkpoint Protein Assays

The plasma levels of 16 soluble immune checkpoint proteins were measured using multiplexed fluorescent bead-based immunoassays with the Milliplex Map Kit (EMD Millipore Corporation, Burlington, MA, USA) and the Luminex Bio-Plex-200 system (Bio-Rad Laboratories, Hercules, CA, USA): namely, sBTLA, sCD27, sCD28, sCD40, sCD80, sCD86, sCTLA-4, sGITR, sGITRL, sHVEM, sPD-1, sPD-L1, sTIM-3, sTLR-2, soluble lymphocyte-activation gene 3 (sLAG-3), and soluble inducible T-cell costimulator (sICOS). This kit enables simultaneous measurement of the concentration of these 16 checkpoint proteins in the plasma sample. In brief, the capture antibody-coupled beads were first incubated with antigen standards or samples for a specific time. The plate was washed to remove unbound materials, followed by incubation with biotinylated detection antibodies. After washing away the unbound biotinylated antibodies, the beads were incubated with a reporter streptavidin–phycoerythrin conjugate (SA–PE). Following the removal of the excess SA–PE, the beads were passed through the array reader, which measures the fluorescence of the bound SA–PE [50]. According to the manufacturer’s instructions, 12.5 µL of plasma was used for each measurement and all samples were assayed in duplicate; mean values were then adopted for further analysis. For values that were lower than the limit of detection, we used 10% of the lowest recorded value as a substitute [51].

4.3. Ethical Considerations

All patients supplied informed consent and the present study was conducted in accordance with the Declaration of Helsinki and was approved by the ethical committee of Osaka City University (#3719, approved on 12 July 2017).

4.4. Statistical Analysis

Analysis was conducted in Easy R (EZR) [52] (Saitama Medical Center, Jichi Medical University, Saitama, Japan) or R [53] (The R Foundation for Statistical Computing, Vienna, Austria) and figures were produced using the package ggplot2 [54]. Categorical variables were compared using Fisher’s
exact test or the chi-squared test, when appropriate. Continuous variables were tested using the Mann–Whitney U test. Wilcoxon signed-rank tests were chosen to compare changes in soluble immune checkpoint concentrations during the early treatment period. Correlations of fold-changes in the levels of two proteins were determined with Spearman’s rank correlation test. A p-value of less than 0.05 was considered to be statistically significant.

5. Conclusions

The present study investigated changes in 16 soluble immune checkpoint proteins in the plasma of patients with HCC treated by lenvatinib or TACE. In the 24 lenvatinib-treated patients, the plasma level of sCD27 decreased but those of sCD40 and sTIM-3 increased significantly at Week 1, while the level of sCD27 was increased significantly at Weeks 2 to 4. These changes in soluble checkpoint protein levels during lenvatinib treatment were different from those seen during sorafenib treatment in our previous study [21], suggesting the two drugs present different inhibitory profiles against the receptor tyrosine kinases. Meanwhile, in the 22 TACE-treated patients, alongside the levels of sCD27, sCD40, and sTIM-3, those of sHVEM, sTLR-2, sCD80, sCTLA-4, sGITR, sGITRL, and sPD-L1 also increased. However, interpretation of the results among TACE-treated patients is difficult because TACE may be accompanied by some confounding factors, as discussed above. Further study is needed to better understand the immunomodulatory effect of the treatments, which may help future investigators to establish an effective combination immunotherapy.

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Conflicts of Interest: Norifumi Kawada has received lecture fees from Eisai Co. Ltd.

Appendix A

Table A1. The 16 immune checkpoint proteins whose soluble forms were analyzed in this study.

| Immune Checkpoint Proteins | Function |
|----------------------------|----------|
| BTLA | B- and T-lymphocyte attenuator | A ligand of HVEM |
| CD27 | Cluster of differentiation 27 | Interaction with CD70 activates antigen-presenting cells |
| CD28 | Cluster of differentiation 28 | Interaction with CD80 or CD86 provides stimulatory signals required for T-cell activation and survival |
| CD40 | Cluster of differentiation 40 | Interaction with CD40L activates antigen-presenting cells |
| CD80 | Cluster of differentiation 80 | A ligand of CD28 and CTLA-4 |
| CD86 | Cluster of differentiation 86 | A ligand of CD28 and CTLA-4 |
| CTLA-4 | Cytotoxic T-lymphocyte antigen 4 | Interaction with either CD80 or CD86 transmits an inhibitory signal to T-cells |
| GITR | Glucocorticoid-induced tumor necrosis factor receptor | Interaction with GITRL inhibits the activity of Tregs and extends the survival of T-effector cells |
| GITRL | Glucocorticoid-induced tumor necrosis factor receptor ligand | A ligand of GITR |
| HVEM | Herpes virus entry mediator | Interaction with LIGHT stimulates the proliferation of T-cells, while interaction with BTLA negatively regulates T-cell responses |
Table A1. Cont.

| Immune Checkpoint Proteins | Function |
|---------------------------|----------|
| ICOS                      | Inducible T-cell costimulator | Interaction with ICOS-L provides a costimulatory signal for T-cell proliferation and survival |
| LAG-3                     | Lymphocyte-activation gene 3 | Interaction with MHC class II inhibits the activation of T-cells |
| PD-1                      | Programmed cell death domain–1 | Interaction with PD-L1 leads to the inhibition of cytotoxic T-cells |
| PD-L1                     | Programmed death-ligand 1 | A ligand of PD-1 |
| TIM-3                     | T-cell immunoglobulin mucin–3 | Interaction with galectin-9 negatively regulates Th1 function by triggering cell death |
| TLR-2                     | Toll-like receptor 2 | TLR-2 recognizes many bacterial, fungal, and other endogenous substances and plays a role in innate immune responses |

Abbreviations: Tregs, regulatory T-cells; LIGHT, homologous to lymphotoxin, exhibits inducible expression and competes with HSV glycoprotein D for binding to herpesvirus entry mediator, a receptor expressed on T-lymphocytes; MHC, major histocompatibility complex.

Figure A1. Changes in soluble immune checkpoint protein levels after 1 week of lenvatinib treatment. Box plots display the sBTLA, sLAG-3, sPD-1, sPD-L1, sICOS, sCD28, sCD86, sCD80, sTLR-2, sGITR, sGITRL, and sCTLA-4 levels in HCC patients at baseline and Week 1 of treatment. The vertical lengths of the boxes indicate the interquartile ranges and the lines in the boxes suggest the median values, while the error bars show the minimum and maximum values (range). The Wilcoxon signed-rank test was used.
Figure A2. Changes in soluble immune checkpoint protein levels after two to four weeks of lenvatinib treatment. Box plots display the sBTLA, sLAG-3, sPD-1, sPD-L1, sICOS, sCD28, sCD86, sCD80, sTLR-2, sGITR, sGITRL, and sCTLA-4 levels in HCC patients at baseline and Week 2 through 4 of treatment. The vertical lengths of the boxes indicate the interquartile ranges and the lines in the boxes show the median values, while the error bars reveal the minimum and maximum values (range). The Wilcoxon signed-rank test was used.

Figure A3. Changes in soluble immune checkpoint protein levels after 1 week of TACE treatment. Box plots display the sBTLA, sLAG-3, sPD-1, sICOS, sCD28, and sCD86 levels in HCC patients at baseline and Week 1 of treatment. The vertical lengths of the boxes reveal the interquartile ranges and the lines in the boxes show the median values, while the error bars suggest the minimum and maximum values (range). The Wilcoxon signed-rank test was used.
Figure A4. The relationships between fold-changes in soluble immune checkpoint protein levels in plasma at two to four weeks of lenvatinib treatment are seen. (a) A positive correlation was found between the fold-changes of soluble checkpoint receptors and their ligands, including sCTLA-4 with sCD80/sCD86 and sPD-1 with sPD-L1, respectively. (b) The fold-change of sCD27 was positively correlated with those of sCD86, sPD-1, and sPD-L1. Spearman’s rank correlation test was used. A p-value of less than 0.05 was considered to be statistically significant.

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