Aplastic anemia and dental implant rehabilitation: a clinical trial

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The purpose of this study was to investigate implant-supported restoration as a technique for restoring missing teeth in patients with aplastic anemia. Recurrent bleeding from wound sites leads to persistent release of iron in the tissue. Excessive iron in tissue is related to clinical findings, including fibrosis, poor wound healing, and high level of angiogenesis, which are possible etiological factors of reduced osseointegration. A 44-year-old female patient with aplastic anemia was treated with multiple endosseous implants throughout the mandible and in the posterior region of the maxilla. After 14 implants were placed, radiological and clinical parameters were assessed during the follow-up period. Marginal bone did not change significantly during the follow-up period. The fine trabecular bone in intimate contact and enclosing the implant fixture was sufficient for successful osseointegration. None of the 14 implants were associated with complications during the seven-year experimental period. This study suggests that dental implant procedures are a safe and reliable treatment option for restoration of missing dentition in patients with aplastic anemia.

Key words: Aplastic anemia, Dental implants, Rehabilitation

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aplastic anemia, infection and bleeding can be life threatening, and such patients have demonstrated a high mortality rate. Although patients with moderate aplastic anemia might not require treatment, frequent and regular blood cell counting is important. Recent therapeutic advances, such as immunosuppressive therapy with bone marrow transplantation, use of growth factor alone or in conjunction with immunosuppressive therapy after bone marrow transplantation, blood component transfusion, and improved infection control, have increased survival rates\(^{10,11}\) of such patients.

This study evaluated the following parameters; 1) the amount of peri-implant bone resorption, 2) level of osseointegration as measured by the amount of fine trabecular bone in close contact with the fixture, 3) probing depth, and 4) peri-abutment gingival connective tissue contour. The purpose of this study was to demonstrate the effectiveness of implant-supported restoration in aplastic anemic patients by evaluating radiographic and clinical features.

II. Case Report

A 44-year-old woman presented to the Department of Oral and Maxillofacial Surgery of Chonnam National University Hospital (Gwangju, Korea) to evaluate missing teeth in the maxilla and the mandible and possible implant-supported restoration. The patient had been diagnosed with severe aplastic anemia at the Department of Hemato-Oncology of Chonnam National University Hospital in July 1995 and had received immunosuppressive therapy. She has been under regular follow-up and had taken 1 mg of folic acid and 50 mg of pyridoxine (vitamin B6) from May 2004 to June 2005.

A general assessment was performed based on panoramic radiograph and cone-beam computed tomography, and we planned to install 14 endosseous implants. Ten implants were inserted in the mandible in the first stage, and four implants were inserted in the maxilla in the second stage, with elevation of both sinuses and autogenous bone grafts. During the preoperative evaluation, the Department of Hemato-Oncology confirmed that the patient would be able to maintain a greater than 1,000 absolute neutrophil count and have lower chances of postoperative bleeding if she was able to maintain a greater than 80,000 platelet count during the surgery. A complete blood cell count investigation revealed that the patient had an RBC of 3.51 million, a WBC count of 3,000, and a count of 84,000 platelets per milliliter of blood. These results indicated that both extraction of teeth #17, #16, #26, and #27 with poor prognosis and implant surgery could be performed without any special consideration.

After five months of healing, the full mucoperiosteal flap was elevated with a midcrestal incision and two vertical releasing incisions under intravenous sedation with midazolam (0.05 mg/kg) and the local anesthesia lidocaine plus with epinephrine 1:100,000. Ten Osstem US II (Osstem Implant Co., Busan, Korea) implants were installed in the spaces left by teeth #32, #33, #34, #36, #37, #42, #43, #44, #46, and #47 (Fig. 1. A), and then provisional implants were placed in the sockets of teeth #31, #35, #41, #45, (Fig. 1. D) Primary closure of the flap involved periosteal releasing incisions, and 3-0 Vicryl (Ethicon Inc., Cornelia, GA, USA) was used as suture material.

Three months after implant installation, a second surgery was performed with an apically repositioned flap in order to produce keratinized gingiva, which helps maintain a healthier gingival environment. (Fig. 1. B) Four months postoperatively, the provisional prostheses were replaced with the final prostheses. (Fig. 1. C)

We obtained panoramic radiographs using the KODAK 8000C System (Carestream Health Inc., Atlanta, GA, USA) and evaluated all panoramic and periapical radiographs with PiView Star (INFINITT Healthcare Co., Ltd., Seoul, Korea). Marginal bone change was estimated from the border between the implant platform and implant abutment to the low-ermost part of the absorbed marginal bone around the implant fixture, based on radiographs performed immediately after the operation. The magnification ratio on the radiographs was corrected by comparing the actual length of an implant fixture with that on the radiographic image.

After two months of loading on the final mandible prostheses, bilateral sinus elevation was performed with ramal bone graft. The oval-shaped bony windows were removed from the lateral walls of the sinuses, and sinus membranes were carefully elevated, followed by bone grafting. Particulated ramal bone graft material was used in combination with Bio-Oss (Geistlich Pharma AG, Wolhusen, Switzerland), and four Osstem US II implants were simultaneously placed at sites #16, #17, #26, and #27. (Fig. 1. E) Fibrin glue was injected into the bone graft site to enhance the bone healing process. All of the procedures were performed under intravenous sedation and local anesthesia with a vasoconstrictor, and the medications were the same as listed above.

Seven months after implant installation in the sinus elevation site, prosthetic loading was applied, and peri-implant bone change was evaluated in panoramic and periapical radiographs. After one year and three months of prosthetic
loading on the lower jaw, marginal bone change was evaluated around the implants using panoramic and periapical radiographs (Fig. 1. F), which revealed peri-implant bone resorption of 1.37 mm on #16, 1.73 mm on #17, 1.30 mm on #26, 1.11 mm on #27, 1.32 mm on #32, 1.70 mm on #33, 1.89 mm on #34, 1.0 mm on #36, 0.86 mm on #37, 0.50 mm on #42, 0.72 mm on #43, 1.0 mm on #44, 2.52 mm on #46, and 2.12 mm on #47.

Tranexamic acid at a concentration of 500 mg/5 mL was available for perioperative bleeding. However, no severe bleeding occurred intraoperatively or postoperatively, and no postoperative complications were noted.

On follow-up, the marginal bone had not changed significantly one year and three months after prosthetic loading. In periapical radiographs, fine trabecular bone was observed around each fixture, which indicates that osseointegration was successful for all of the implants. We measured the probing depth, which is the distance from the gingival margin to the periodontal pocket, on the mesiobuccal and distobuccal sides of each tooth. The average of the measured values for each tooth was calculated, and a mean probing depth of 1.66 mm was obtained using the Williams Colorvue ProbeKit.
bone resorption, observed while loading dental prostheses, other implants were well-maintained, fulfilling the success criteria cited by Albrektsson et al. The first two implants were categorized into group II (satisfactory survival), and the others into group I (success, optimum health), according to the health scale developed by the International Congress of Oral Implantologists Consensus Conference for Implant Success in Pisa in 2007. Fine trabecular bone structure was observed around all of the implant fixtures, which created sufficient anchorage necessary for long-term implant maintenance. Peri-implant resorption did not change significantly during the follow-up period. Stable, rigid, fixated implants have been reported with pocket depths ranging from 2 to 6 mm; in the present case, the mean probing depth was 1.66 mm after 21.4 months of prosthetic loading. There were no complications during the seven-year follow-up period. Thus, based on these results, implant surgery can be performed safely even for patients with hematologic disorders by applying three principles of implant installation; Induction of initial hemostasis, maintenance of hemostasis during the healing period, and immobilization of the implant. These can be achieved through appropriate systemic treatment and precise management of local soft tissue.

III. Discussion

Implant-supported restoration has become common practice in dentistry and is associated with long-term successful outcomes. This could increase the popularity of implant-supported restoration among dentists and the general public even though its effect in medically compromised patients, especially those with bleeding disorders, has not been thoroughly demonstrated.

Special considerations for these medically compromised patients include bleeding control during operative and post-operative periods in order to avoid life-threatening situations, achieving successful osseointegration, and long-term implant maintenance. Restoration of initial hemostasis can modulate some of the parameters of wound healing. An extended period of adequate hemostatic function is necessary to accommodate the normal healing process, probably because the risk of hemorrhage is increased by vascular remodeling and angiogenesis during the healing process. Furthermore, patients with aplastic anemia are susceptible to peri-implant infection as aplastic anemia is characterized by marked hypoplastic bone marrow and pancytopenia (anemia, leucopenia, and thrombocytopenia).

We were not able to collect data regarding specific blood management strategies for successful implant placement in cases of aplastic anemia, so we followed the general guidelines. We prepared a blood transfusion to maintain platelet count during surgery, since a platelet count less than 25,000/mL indicates a high risk of spontaneous oral bleeding. Tranexamic acid was also prepared for prolonged retention of intact weak blood clots and to prevent bleeding in plasmin-rich areas such as the oral cavity. Tranexamic acid can significantly reduce blood loss after oral surgery in patients with hemophilia and can be used topically or systemically.

Radiographic examinations with implant mobility tests are the most reliable for assessing osseointegration. Although two of the present implants showed moderate peri-implant bone resorption, observed while loading dental prostheses, other implants were well-maintained, fulfilling the success criteria cited by Albrektsson et al. The first two implants were categorized into group II (satisfactory survival), and the others into group I (success, optimum health), according to the health scale developed by the International Congress of Oral Implantologists Consensus Conference for Implant Success in Pisa in 2007. Fine trabecular bone structure was observed around all of the implant fixtures, which created sufficient anchorage necessary for long-term implant maintenance. Peri-implant resorption did not change significantly during the follow-up period. Stable, rigid, fixated implants have been reported with pocket depths ranging from 2 to 6 mm; in the presented case, the mean probing depth was 1.66 mm after 21.4 months of prosthetic loading. There were no complications during the seven-year follow-up period.

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Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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