

# VCA - head and neck transplant recipient registration



OMB No. 0915-0157, Expiration date: 9/30/2026

The transplant recipient registration (TRR) forms are generated and available after a transplant event is reported to the OPTN. The TRR record is completed by the transplant hospital performing the transplant. The registration and hospital discharge follow-up information is combined in this record.

Complete the TRR at hospital discharge or six weeks post-transplant, whichever is first. If the recipient is still hospitalized at six weeks post-transplant, provide the most recent information available regarding the recipient's progress.

Complete one TRR form for recipients of bilateral upper limbs. Complete separate TRR forms for each VCA organ transplant.

The TRR must be validated within 90 days of the record generation date. Example: If the recipient is removed as being transplanted on 10/1/XXXX, the TRR form will be due 90 days from that date, 12/30/XXXX. See OPTN Policies (<https://optn.transplant.hrsa.gov/policies-bylaws/policies>) for additional information.

## Recipient information

| Question                                | Answers  |
|---|--|
| Surgical procedure (prepopulated)*      | <input type="checkbox"/> Face<br><input type="checkbox"/> Larynx<br><input type="checkbox"/> Scalp<br><input type="checkbox"/> Trachea<br><input type="checkbox"/> Vascularized parathyroid gland<br><input type="checkbox"/> Vascularized thyroid |
| Recipient first name (prepopulated)*    | _____  |
| Recipient last name (prepopulated)*     | _____  |
| Recipient middle initial (prepopulated) | _____  |
| Date of birth (prepopulated)*           | _____  |
| SSN (prepopulated)*                     | _____  |
| Birth sex (prepopulated)*               | _____  |
| HIC                                     | _____  |
| Transplant date (prepopulated)*         | _____  |
| State of permanent residence*           | _____  |
| Permanent ZIP code                      | _____  |
| Expected date (prepopulated)*           | _____  |

## Provider information

| Question                           | Answers |
|------------------------------------|---------|
| Recipient center (prepopulated)*   | _____   |
| Lead reconstructive surgeon name*  | _____   |
| Lead reconstructive surgeon NPI #* | _____   |

## Donor information

| Question                        | Answers |
|---------------------------------|---------|
| UNOS donor ID # (prepopulated)* | _____   |

Donor type (prepopulated)\* \_\_\_\_\_

OPO (prepopulated) \_\_\_\_\_

## Patient status - Transplant hospitalization

| Question                                | Answers |
|---|---------|
| Date of admission to transplant center* | _____   |
| Date of discharge from hospital         | _____   |

## Patient status

| Question  | Answers |
|---|---------|
| Date last seen, graft removed, or death*                                | _____   |
| Patient status*   | _____   |
| <i>If patient status is "Dead", select the patient's cause of death</i> |         |
| Primary cause of death  | _____   |
| Other, specify  | _____   |

## Socio-demographic information: Pre-transplant

| Question                                       | Answers |
|--|---------|
| Highest education level*                       | _____   |
| <i>For recipients 18 years of age or older</i> |         |
| Working for income                             | _____   |

## Socio-demographic information: Pre-transplant - Source of payment

| Question   | Answers |
|--|---------|
| Grant funding*   | _____   |
| Institutional funding*   | _____   |
| Primary source of payment*   | _____   |
| <i>If primary source of payment is "Foreign government, specify" select foreign government</i> |         |
| Primary source of payment - foreign government, specify  | _____   |

## Clinical information: Pre-transplant

| Question | Answers |
|----------|---------|
|----------|---------|

Enter height or height status

Height (cm: 1.00-225.00)

Height status

Enter weight or weight status

Weight (kg: 0.45-294.84)

Weight status

Primary diagnosis for transplant\*

Primary diagnosis for transplant - other, specify

Previous transplants (VCA or non-VCA organs)\*

Previous skin graft(s)\*

Was patient hospitalized during the last 90 days prior to the transplant admission\*

Medical condition at time of transplant\*

Any tolerance induction technique used\*

Pre-transplant blood transfusions\*

For recipients whose birth sex is female

Number of pre-transplant pregnancies (which may or may not have resulted in a live birth: 0-50)

Malignancies prior to transplant\*

If malignancies prior to transplant is "Yes", select type

Specify type (select all that apply)

- Breast
- CNS tumor
- Genitourinary
- Hepatocellular carcinoma
- Leukemia/Lymphoma
- Liver
- Lung
- Skin: melanoma
- Skin: non-melanoma
- Thyroid
- Tongue/throat/larynx
- Other, specify

Other, specify

## Clinical information: Pre-transplant - Amount of tissue loss

Question

Craniofacial

If "Partial", specify anatomic structures missing

Answers

If "Other specify", specify amount of tissue loss

## Clinical information: Pre-transplant - Viral detection

| Question   | Answers |
|--|---------|
| HIV serostatus*  | _____   |
| CMV status*  | _____   |
| HBV core antibody*   | _____   |
| HBV surface antigen*   | _____   |
| HCV serostatus*  | _____   |
| EBV serostatus*  | _____   |
| Did the recipient receive Hepatitis B vaccines prior to transplant?* | _____   |
| Reason not vaccinated  | _____   |
| Other, specify   | _____   |

## Clinical information: Pre-transplant - Pre-transplant labs

| Question   | Answers |
|--|---------|
| <i>Enter serum creatinine or serum creatinine status</i> |         |
| Serum creatinine (mg/dL: 0.10-25.00)                     | _____   |
| Serum creatinine status                                  | _____   |
| <i>Enter hemoglobin A1c or hemoglobin A1c status</i>     |         |
| Hemoglobin A1c (%: 0-100)                                | _____   |
| Hemoglobin A1c status                                    | _____   |
| Calculated PRA (CPRA) at transplant (%: 0-100)*          | _____   |
| Donor crossmatch result*                                 | _____   |

## Functional status: Pre-transplant

| Question   | Answers |
|--|---------|
| <i>For recipients younger than 18 years of age at transplant</i> |         |
| Motor development  | _____   |

## Functional status: Pre-transplant - SF-12 score: Physical health

| Question                                    | Answers |
|---|---------|
| Physical functioning (PF) score (0.0-100.0) | _____   |
| Role-physical (RP) score (0.0-100.0)        | _____   |
| Bodily pain (BP) score (0.0-100.0)          | _____   |
| General health (GH) score (0.0-100.0)       | _____   |

Physical component summary (PCS) score (0.0-100.0)

\_\_\_\_\_

## Functional status: Pre-transplant - SF-12 score: Mental health

**Question**

**Answers**

Vitality (VT) score (0.0-100.0)

\_\_\_\_\_

Social functioning (SF) score (0.0-100.0)

\_\_\_\_\_

Role-emotional (RE) score (0.0-100.0)

\_\_\_\_\_

Mental health (MH) score (0.0-100.0)

\_\_\_\_\_

Mental component summary (MCS) score (0.0-100.0)

\_\_\_\_\_

## Clinical information: Transplant procedure

**Question**

**Answers**

Multiple graft recipient\*

\_\_\_\_\_

Were extra allograft vessels/nerve/tissue from outside the donated graft used in the transplant procedure\*

\_\_\_\_\_

## Clinical information: Transplant procedure - Preservation information

**Question**

**Answers**

Warm ischemia time (include anastomotic time; minutes: 0-2880)

\_\_\_\_\_

Cold ischemia time (minutes: 0-2880)

\_\_\_\_\_

## Clinical information: Post-transplant

**Question**

**Answers**

Graft status\*

\_\_\_\_\_

*If "Failed"*

Date of graft failure

\_\_\_\_\_

*Causes of graft failure*

Acute rejection

\_\_\_\_\_

Acute rejection - Banff score

\_\_\_\_\_

Acute rejection - visual skin changes

\_\_\_\_\_

Chronic rejection

\_\_\_\_\_

Chronic rejection - visual skin changes

\_\_\_\_\_

Vascular complications

\_\_\_\_\_

Sepsis / Infection

\_\_\_\_\_

Trauma

\_\_\_\_\_

|   |       |
|---|-------|
| Patient requested removal   | _____ |
| Non-adherence   | _____ |
| Other   | _____ |
| Other, specify  | _____ |
| _____   | _____ |
| Did patient have any acute rejection episodes between transplant and discharge* | _____ |
| If yes, number of rejection episodes (1-100)                                    | _____ |
| <i>Enter for each episode</i>   |       |
| Date of acute rejection diagnosis   | _____ |
| Acute rejection was treated   | _____ |
| Visual skin changes   | _____ |
| Biopsy was done to confirm acute rejection                                      | _____ |
| Banff score   | _____ |
| <i>Enter for each episode</i>   |       |
| Date of acute rejection diagnosis   | _____ |
| Acute rejection was treated   | _____ |
| Visual skin changes   | _____ |
| Biopsy was done to confirm acute rejection                                      | _____ |
| Banff score   | _____ |
| <i>Enter for each episode</i>   |       |
| Date of acute rejection diagnosis   | _____ |
| Acute rejection was treated   | _____ |
| Visual skin changes   | _____ |
| Biopsy was done to confirm acute rejection                                      | _____ |
| Banff score   | _____ |
| <i>Enter for each episode</i>   |       |
| Date of acute rejection diagnosis   | _____ |
| Acute rejection was treated   | _____ |
| Visual skin changes   | _____ |
| Biopsy was done to confirm acute rejection                                      | _____ |
| Banff score   | _____ |
| <i>Enter for each episode</i>   |       |
| Date of acute rejection diagnosis   | _____ |
| Acute rejection was treated   | _____ |
| Visual skin changes   | _____ |
| Biopsy was done to confirm acute rejection                                      | _____ |
| Banff score   | _____ |

**Clinical information: Post-transplant - Lab data at time of discharge from the hospital**

| Question | Answers |
|----------|---------|
|----------|---------|

Enter serum creatinine or serum creatinine status

Serum creatinine (mg/dL: 0.10-25.00)

\_\_\_\_\_

Serum creatinine status

\_\_\_\_\_

Enter hemoglobin A1c or hemoglobin A1c status

Hemoglobin A1c (%: 0-100)

\_\_\_\_\_

Hemoglobin A1c status

\_\_\_\_\_

## Clinical information: Post-transplant - Major transplant complication

**Question**

**Answers**

Arterial thrombosis\*

\_\_\_\_\_

Venous thrombosis\*

\_\_\_\_\_

More than 5 pRBC (packed red blood cells) units\*

\_\_\_\_\_

Cardiac arrest\*

\_\_\_\_\_

DIC (Disseminated intravascular coagulation)\*

\_\_\_\_\_

Graft/reperfusion syndrome\*

\_\_\_\_\_

Other major transplant complications

\_\_\_\_\_

Other major transplant complications - other, specify

\_\_\_\_\_

## Functional status: Post-transplant - Head and neck

**Question**

**Answers**

Smile restoration

\_\_\_\_\_

Ability to open and close eyelids

\_\_\_\_\_

## Treatment

**Question**

**Answers**

Antiviral prophylaxis\*

\_\_\_\_\_

Antibacterial prophylaxis\*

\_\_\_\_\_

Antifungal prophylaxis\*

\_\_\_\_\_

Peri-operative anticoagulation\*

\_\_\_\_\_

## Topical immunosuppressive medication - Topical drugs used for acute rejection or maintenance

**Question**

**Answers**

\_\_\_\_\_

|   |   |
|---|---|
| Steroids (Clobetasol)                             | <input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |
| Tacrolimus (Protopic)                             | <input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |
| Other, specify 1                                  | _____   |
| Other, specify 1 - Acute rejection or maintenance | <input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |
| Other, specify 2                                  | _____   |
| Other, specify 2 - Acute rejection or maintenance | <input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |

## Non-topical immunosuppressive medication - Drugs used for induction, acute rejection, or maintenance

| Question  | Answers  |
|---|--|
| Steroids (prednisone, methylprednisolone, Solumedrol, Medrol, Decadron)                                       | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |
| Steroids (prednisone, methylprednisolone, Solumedrol, Medrol, Decadron) - Number of days of induction (0-365) | _____  |
| Steroids (prednisone, methylprednisolone, Solumedrol, Medrol, Decadron) - Status                              | _____  |

## Non-topical immunosuppressive medication - Drugs used for induction or acute rejection

| Question   | Answers   |
|--|---|
| Atgam  | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Anti-rejection indication |
| Atgam - Number of days of induction (0-365)                            | _____   |
| Atgam - Status   | _____   |
| Campath (alemtuzumab, anti-CD52)                                       | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Anti-rejection indication |
| Campath (alemtuzumab, anti-CD52) - Number of days of induction (0-365) | _____   |
| Campath (alemtuzumab, anti-CD52) - Status                              | _____   |
| Cytoxan (cyclophosphamide)   | <input type="checkbox"/> Anti-rejection indication  |
| Methotrexate (Folex PFS, Mexate-AQ, Rheumatrex)                        | <input type="checkbox"/> Anti-rejection indication  |
| OKT3 (Orthoclone, muromonab)   | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Anti-rejection indication |
| OKT3 (Orthoclone, muromonab) - Number of days of induction (0-365)     | _____   |
| OKT3 (Orthoclone, muromonab) - Status                                  | _____   |
| Rituxan (rituximab)  | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Anti-rejection indication |
| Rituxan (rituximab) - Number of days of induction (0-365)              | _____   |



|  |   |
|--|---|
| Rituxan (rituximab) - Status                                 | _____   |
| Simulect (basiliximab)                                       | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Anti-rejection indication |
| Simulect (basiliximab) - Number of days of induction (0-365) | _____   |
| Simulect (basiliximab) - Status                              | _____   |
| Thymoglobulin  | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Anti-rejection indication |
| Thymoglobulin - Number of days of induction (0-365)          | _____   |
| Thymoglobulin - Status                                       | _____   |

## Non-topical immunosuppressive medication - Drugs primarily used for maintenance

| Question  | Answers   |
|---|---|
| <i>Cyclosporine, select from the following:</i>                       |   |
| EON (generic cyclosporine)  | <input type="checkbox"/> Maintenance indicator  |
| Gengraf (Abbott cyclosporine)   | <input type="checkbox"/> Maintenance indicator  |
| Neoral (CyA-NOF)  | <input type="checkbox"/> Maintenance indicator  |
| Other generic cyclosporine, specify brand                             | _____   |
| Other generic cyclosporine  | <input type="checkbox"/> Maintenance indicator  |
| Sandimmune (cyclosporine A)   | <input type="checkbox"/> Maintenance indicator  |
| Imuran (azathioprine, AZA)  | <input type="checkbox"/> Maintenance indicator  |
| Leflunomide (LFL)   | <input type="checkbox"/> Maintenance indicator  |
| <i>Mycophenolate acid, select from the following:</i>                 |   |
| CellCept (MMF)  | <input type="checkbox"/> Maintenance indicator  |
| Generic MMF (generic CellCept)  | <input type="checkbox"/> Maintenance indicator  |
| Myfortic (mycophenolate acid)   | <input type="checkbox"/> Maintenance indicator  |
| Nulojix (belatacept)  | <input type="checkbox"/> Maintenance indicator  |
| Rapamune (sirolimus, Rapamycin)                                       | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Maintenance indicator |
| Rapamune (sirolimus, Rapamycin) - Number of days of induction (0-365) | _____   |
| Rapamune (sirolimus, Rapamycin) - Status                              | _____   |
| <i>Tacrolimus, select from the following:</i>                         |   |
| Astagraf XL (extended release tacrolimus)                             | <input type="checkbox"/> Maintenance indicator  |
| Generic tacrolimus (generic Prograf)                                  | <input type="checkbox"/> Maintenance indicator  |
| Prograf (FK506)   | <input type="checkbox"/> Maintenance indicator  |
| Zortress (everolimus)   | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Maintenance indicator |

|  |  |
|--|--|
| Zortress (everolimus) - Number of days of induction (0-365)                | _____  |
| Zortress (everolimus) - Status   | _____  |
| <b>Other drugs</b>   |  |
| Other immunosuppressive medication, specify:                               | _____  |
| Other immunosuppressive medication 1                                       | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |
| Other immunosuppressive medication 1 - Number of days of induction (0-365) | _____  |
| Other immunosuppressive medication 1 - Status                              | _____  |
| Other immunosuppressive medication, specify:                               | _____  |
| Other immunosuppressive medication 2                                       | <input type="checkbox"/> Induction indication<br><input type="checkbox"/> Maintenance indication<br><input type="checkbox"/> Anti-rejection indication |
| Other immunosuppressive medication 2 - Number of days of induction (0-365) | _____  |
| Other immunosuppressive medication 2 - Status                              | _____  |

## Public Burden/Privacy Act Statements

Department of Health and Human Services  
Health Resources and Services Administration

OMB No: 0915-0157  
Expiration Date: 9/30/2026

# ORGAN PROCUREMENT AND TRANSPLANTATION NETWORK (OPTN)

## DATA COLLECTION

**DATA ACCURACY CERTIFICATION:** I certify that the data entered by me in UNet<sup>SM</sup> are accurate, timely, and complete to the best of my knowledge, information and belief. These data are based upon information contained in corresponding medical records and other source documents, or where appropriate, are based upon clinical observation.

**PUBLIC BURDEN STATEMENT:** An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. The OMB control number for this project is 0915-0157. Public reporting burden for the applicant for this collection of information is estimated to average 53 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to HRSA Reports Clearance Officer, 5600 Fishers Lane, Room 1033, Rockville, Maryland 20857.

**PRIVACY ACT STATEMENT:** In accordance with the requirements of the Privacy Act of 1974 (<https://www.federalregister.gov/documents/2022/08/01/2022-16344/privacy-act-of-1974-system-of-records>) as amended, 42 U.S.C. § 273, et seq., and 42 CFR Part 121 authorize collection of this information by the OPTN. This information is distributed to the Scientific Registry of Transplant Recipients (SRTR) and the Health Resources and Services Administration (HRSA), with the United States Department of Health and Human Services. The primary uses of this information are to match organ donors with recipients, to monitor compliance of member organizations with OPTN requirements, to review and report on the status of organ donation and transplantation in the United States, and to provide data to researchers and government agencies to study transplantation. The routine uses which may be made of this information are: (i) to organ procurement organizations and transplant hospitals to match organ donors with compatible recipients and validate the accuracy of donor and recipient; (ii) to the Department of Justice to use in defending litigation; (iii) to a congressional office upon the request of an individual concerning records pertaining to him/her; (iv) for research purposes, if certain requirements are satisfied and data use agreements are executed; and (v) to Agency contractors who have been engaged by the Agency to assist in accomplishment of an Agency function relating to the purposes of this system and who need to have access to the records in order to assist the Agency. Furnishing the remaining information requested is required by law of organ procurement organizations and transplant hospitals and the failure to submit such information may result in enforcement actions resulting from noncompliance with OPTN requirements.  
HRSA (08/02)

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