Section 3.11 General and Informed Consent to Treatment

3.11.1 Introduction
Each behavioral health recipient has the right to participate in decisions regarding his or her behavioral health care, including the right to refuse treatment. It is important for persons seeking behavioral health services to agree to those services and be made aware of the service options and alternatives available to them as well as specific risks and benefits associated with these services.

The Arizona Department of Health Services/Division of Behavioral Health Services (ADHS/DBHS) recognizes two primary types of consent: general consent and informed consent.

General consent is a one-time agreement to receive behavioral health services that is usually obtained from a person during the intake process at the initial appointment, and is always obtained prior to the provision of any behavioral health services. General consent must be verified by a behavioral health recipient’s or legal guardian’s signature.

Informed consent must be obtained before the provision of a specific treatment that has associated risks and benefits. Informed consent is required prior to the provision of the following services and procedures:

- Complementary and Alternative Medicine (CAM);
- Psychotropic medications;
- Electro-convulsive therapy (ECT);
- Use of telemedicine;
- Application for a voluntary evaluation;
- Research;
- Admission for medical detoxification, an inpatient facility or a residential program (for persons determined to have a Serious Mental Illness); and
- Procedures or services with known substantial risks or side effects

Prior to obtaining informed consent, an appropriate behavioral health representative, as identified in R9-21-206.01(c), must present the facts necessary for a person to make an informed decision.
informed decision regarding whether to agree to the specific treatment and/or procedures. Documentation that the required information was given and that the person agrees or does not agree to the specific treatment must be included in the comprehensive clinical record, as well as the person’s/guardian’s signature when required.

In addition to general and informed consent for treatment, state statute (A.R.S. § 15-104) requires written consent from a child’s parent or legal guardian for any behavioral health survey, analysis, or evaluation conducted in reference to a school based prevention program. (See subsection 3.11.3-E)

The intent of this section is to describe the requirements for reviewing and obtaining general and informed consent, for persons receiving services within the public behavioral health system, as well as consent for any behavioral health survey or evaluation in connection with an ADHS/DBHS school-based prevention program.

3.11.2 Terms
Definitions for terms are located online at www.narbha.org or http://www.azdhs.gov/bhs/definitions/index.php [T/RBHA to insert T/RBHA link]. The following terms are referenced in this section:

Behavioral Health Medical Practitioner

Behavioral Health Professional

General Consent

Informed Consent

Telemedicine

Voluntary Evaluation

3.11.3 Procedures
3.11.3-A. General Requirements
Any person, aged 18 years and older, in need of behavioral health services must give voluntary general consent to treatment, demonstrated by the person’s or legal guardian’s signature on a general consent form, before receiving behavioral health services.

For persons under the age of 18, the parent, legal guardian, or a lawfully authorized custodial agency must give general consent to treatment, demonstrated by the parent, legal guardian, or a lawfully authorized custodial agency representative’s signature on a general consent form prior to the delivery of behavioral health services.

Any person aged 18 years and older or the person’s legal guardian, or in the case of persons under the age of 18, the parent, legal guardian or a lawfully authorized custodial agency, after being fully informed of the consequences, benefits and risks of treatment, has the right not to consent to receive behavioral health services.
Any person aged 18 years and older or the person’s legal guardian, or in the case of persons under the age of 18, the parent, legal guardian or a lawfully authorized custodial agency has the right to refuse medications unless specifically required by a court order or in an emergency situation.

Providers treating persons in an emergency situation are not required to obtain general consent prior to the provision of emergency services. Providers treating persons pursuant to court order must obtain consent, as applicable, in accordance with A.R.S. Title 36, Chapter 5.

All evidence of informed consent and general consent to treatment must be documented in the comprehensive clinical record per Section 4.2, Behavioral Health Medical Record Standards.

3.11.3-B. General Consent
Administrative functions associated with a behavioral health recipient’s enrollment do not require consent, but before any services are provided, general consent must be obtained. General consent is usually obtained during the intake process and represents a person’s, or if under the age of 18, the person’s parent, legal guardian or lawfully authorized custodial agency representative’s, written agreement to participate in and to receive non-specified (general) behavioral health services.

3.11.3-C. Informed Consent
What Information must be provided to obtain informed consent?
In all cases where informed consent is required by this policy, informed consent must include at a minimum:

- Behavioral health recipient’s right to participate in decisions regarding his or her health care, including the right to refuse treatment, and to express preferences about future treatment decisions;

- Information about the person’s diagnosis and the proposed treatment, including the intended outcome, nature and all available procedures involved in the proposed treatment;

- The risks, including any side effects, of the proposed treatment, as well as the risks of not proceeding;

- The alternatives to the proposed treatment, particularly alternatives offering less risk or other adverse effects;

- That any consent given may be withheld or withdrawn in writing or orally at any time. When this occurs the provider must document the person’s choice in the medical record;

- The potential consequences of revoking the informed consent to treatment; and

- A description of any clinical indications that might require suspension or termination of the proposed treatment.
For persons under 18 years of age, the FDA status of the medication and the level of evidence supporting the recommended medication.

Who can give informed consent, and how is it documented?
Persons, or if applicable the client’s parent, guardian or custodian shall give informed consent for treatment by signing and dating an acknowledgment that he or she has received the information and gives informed consent to the proposed treatment.

When informed consent is given by a third party, the identity of the third party and the legal capability to provide consent on behalf of the person, must be established. If the informed consent is for psychotropic medication or telemedicine and the person, or if applicable, the person’s guardian refuses to sign an acknowledgment and gives verbal informed consent, the medical practitioner shall document in the person’s record that the information was given, the client refused to sign an acknowledgment and that the client gives informed consent to use psychotropic medication or telemedicine.

Who can provide informed consent and how is it communicated?
When providing information that forms the basis of an informed consent decision for the circumstances identified above, the information must be:

- Presented in a manner that is understandable and culturally appropriate to the person, parent, legal guardian or an appropriate court; and
- Presented by a credentialed behavioral health medical practitioner or a registered nurse with at least one year of behavioral health experience. It is preferred that the prescribing clinician provide information forming the basis of an informed consent decision. In a specific situation in which that is not possible or practicable, information may be provided by another credentialed behavioral health medical practitioner or registered nurse with at least one year of behavioral health experience.

Psychotropic Medications, Complementary and Alternative Treatment and Telemedicine
Unless treatments and procedures are court ordered, providers must obtain written informed consent, and if written consent is not obtainable, providers must obtain oral informed consent. If oral informed consent is obtained instead of written consent from the person, parent or legal guardian, it must be documented in written fashion. Informed consent is required in the following circumstances:

- Prior to the initiation of any psychotropic medication or initiation of Complementary and Alternative Treatment (CAM) (see Section 3.15, Psychotropic Medication Prescribing and Monitoring). The use of PM Form 3.15.1 is recommended as a tool to review and document informed consent for psychotropic medications; and

- Prior to the delivery of behavioral health services through telemedicine. (PM Form 3.11.2)

Electro-Convulsive Therapy (ECT), research activities, voluntary evaluation and procedures or services with known substantial risks or side effects.
Written informed consent must be obtained from the person, parent or legal guardian, unless treatments and procedures are under court order, in the following circumstances:

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Effective Date: 08/01/2014
• Before the provision of (ECT);

• Prior to the involvement of the person in research activities;

• Prior to the provision of a voluntary evaluation for a person. The use of ADHS/DBHS Form MH-103 is required for persons determined to have a Serious Mental Illness and is recommended as a tool to review and document informed consent for voluntary evaluation of all other populations; and

• Prior to the delivery of any other procedure or service with known substantial risks or side effects.

Additional Provisions
Written informed consent must be obtained from the person, legal guardian or an appropriate court prior to the person’s admission to any medical detoxification, inpatient facility or residential program operated by a behavioral health provider.

Revocation of Informed Consent
If informed consent is revoked, treatment must be promptly discontinued, except in cases in which abrupt discontinuation of treatment may pose an imminent risk to the person. In such cases, treatment may be phased out to avoid any harmful effects.

3.11.3-D. Special requirements for children
In accordance with A.R.S. § 36-2272, except as otherwise provided by law or a court order, no person, corporation, association, organization or state-supported institution, or any individual employed by any of these entities, may procure, solicit to perform, arrange for the performance of or perform mental health screening in a nonclinical setting or mental health treatment on a minor without first obtaining consent of a parent or a legal custodian of the minor child. If the parental consent is given through telemedicine, the health professional must verify the parent’s identity at the site where the consent is given. This section does not apply when an emergency exists that requires a person to perform mental health screening or provide mental health treatment to prevent serious injury to or save the life of a minor child.

Transitioning Youth and Psychotropic Medications
Youth under the age of 18 are educated on options, allowed to provide input, and encouraged to assent to medication(s) being prescribed. Information is discussed with the youth in a clear and age-appropriate manner consistent with the developmental needs of the youth. Discussion of the youth’s ability to give consent for medications at the age of 18 years old is begun no later than age 17 ½ years old. There should be special attention to the effect of medications on reproductive status and pregnancy, as well as long term effects on weight, abnormal involuntary movements and other health parameters. Evidence of the youth’s consent to continue medications after his/her 18th birthday may be documented through use of PM Form 3.15.1, a recommended tool to review and document informed consent for psychotropic medications.

Non-emergency Situations
In cases where the parent is unavailable to provide general or informed consent and the child is being supervised by a caregiver who is not the child’s legal guardian (e.g., grandparent) and
does not have power of attorney, general and informed consent must be obtained from one of the following:

- Lawfully authorized legal guardian;

- Foster parent, group home staff or other person with whom the Department of Economic Security/Child Protective Services (DES/CPS) has placed the child; or

- Government agency authorized by the court.

If someone other than the child’s parent intends to provide general and, when applicable, informed consent to treatment, the following documentation must be obtained and filed in the child’s comprehensive clinical record:

<table>
<thead>
<tr>
<th>Individual/Entity</th>
<th>Documentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Legal guardian</td>
<td>Copy of court order assigning custody</td>
</tr>
<tr>
<td>Relatives</td>
<td>Copy of power of attorney document</td>
</tr>
<tr>
<td>Other person/agency</td>
<td>Copy of court order assigning custody</td>
</tr>
<tr>
<td>DES/CPS Placements (for children removed from the home by DES/DPS), such as:</td>
<td>None required</td>
</tr>
<tr>
<td>Foster parents</td>
<td>If behavioral health providers doubt whether the individual bringing the child in for services is a person/agency representative in whose care DES/CPS has placed the child, the provider may ask to review verification, such as documentation given to the individual by DES indicating that the individual is an authorized DES/CPS placement. If the individual does not have this documentation, then the provider may also contact the child's DES/CPS caseworker to verify the individual's identity.</td>
</tr>
<tr>
<td>Group home staff</td>
<td></td>
</tr>
<tr>
<td>Foster home staff</td>
<td></td>
</tr>
<tr>
<td>Relatives</td>
<td></td>
</tr>
<tr>
<td>Other person/agency in whose care</td>
<td></td>
</tr>
<tr>
<td>DES/CPS has placed the child</td>
<td></td>
</tr>
</tbody>
</table>

For any child who has been removed from the home by CPS, the foster parent, group home staff, foster home staff, relative or other person or agency in whose care the child is currently placed may give consent for the following behavioral health services:

- Evaluation and treatment for emergency conditions that are not life threatening; and

- Routine medical and dental treatment and procedures, including early periodic screening, diagnosis and treatment services, and services by health care providers to relieve pain or treat symptoms of common childhood illnesses or conditions (including behavioral health services and psychotropic medications).

**Youth Assent for Psychotropic Medications**

In addition to requirements for all transitioning youth on psychotropic medications, youth, who are not in the custody of their parents, need special attention so that the youth do not have a lapse in care upon turning 18yo. The youth should be active participants in care, understand the reasons for psychotropic medication treatment, give ongoing assent and be able to give informed consent as part of transitioning to adulthood successfully.
Any minor who has entered into a lawful contract of marriage, whether or not that marriage has been dissolved subsequently, emancipated youth or any homeless minor may provide general and, when applicable, informed consent to treatment without parental consent (A.R.S. § 44-132).

Emergency Situations
In emergency situations involving a child in need of immediate hospitalization or medical attention, general and, when applicable, informed consent to treatment is not required.

Any child, 12 years of age or older, who is determined upon diagnosis of a licensed physician, to be under the influence of a dangerous drug or narcotic, not including alcohol, may be considered an emergency situation and can receive behavioral health care as needed for the treatment of the condition without general and, when applicable, informed consent to treatment.

3.11.3-E. Informed Consent during Involuntary Treatment
At times, involuntary treatment can be necessary to protect safety and meet needs when a person, due to mental disorder, is unwilling or unable to consent to necessary treatment. In this case, a court order may serve as the legal basis to proceed with treatment. However, capacity to give informed consent is situational, not global, as an individual may be willing and able to give informed consent for aspects of treatment even when not able to give general consent. Individuals should be assessed for capacity to give informed consent for specific treatment and such consent obtained if the individual is willing and able, even though the individual remains under court order.

3.11.3-F Consent for behavioral health survey or evaluation for school-based prevention programs
Written consent must be obtained from a child’s parent or legal guardian for any behavioral health survey, analysis or evaluation conducted in reference to a school-based prevention program administered by ADHS/DBHS.

Provider Manual Form 3.11.1, Substance Abuse Prevention Program and Evaluation Consent must be used to gain parental consent for evaluation of school based prevention programs. Providers may use an alternative consent form only with the prior written approval of ADHS/DBHS. The consent must satisfy all of the following requirements:

- Contain language that clearly explains the nature of the screening program and when and where the screening will take place;

- Be signed by the child’s parent or legal guardian; and

- Provide notice that a copy of the actual survey, analysis or evaluation questions to be asked of the student is available for inspection upon request by the parent or legal guardian.

Completion of Provider Manual Form 3.11.1, Substance Abuse Prevention Program and Evaluation Consent applies solely to consent for a survey, analysis, or evaluation only, and does not constitute consent for participation in the program itself.
3.11.4 References
The following citations can serve as additional resources for this content area:

20 U.S.C. § 1232h (b)
42 C.F.R. § 438.100
42 C.F.R. § 438.102
A.R.S. § 8-514.05
A.R.S. § 14-5104
A.R.S. § 15-104
A.R.S. § 36-522
A.R.S. § 36-501.21
A.R.S. § 36-2272
A.R.S. § 44-132
R9-20-203
R9-20-208
R9-21-206
R9-21-503
AHCCCS/ADHS Contract
ADHS/RBHA Contracts
ADHS/TRBHA IGAs
ADHS/DBHS Covered Behavioral Health Services Guide
ADHS/DBHS Policy and Procedure Manual CO 1.3, Use of Telemedicine
Section 3.15, Psychotropic Medications: Prescribing and Monitoring
Section 3.18, Pre-Petition Screening, Court-Ordered Evaluation, and Court-Ordered Treatment
Section 4.1 Disclosure of Behavioral Health Information
Section 4.2, Behavioral Health Medical Record Standards
ADHS/DBHS Practice Protocol, Comprehensive Assessment and Treatment for Substance Use Disorders in Children and Adolescents (Formerly known as Practice Improvement Protocol # 10)
The Arizona Medical Board’s Guidelines for Physicians Who Incorporate or Use Complementary or Alternative Medicine in Their Practice

3.11.5 PM Forms

PM Form 3.11.1, Substance Abuse Prevention Program and Evaluation Consent
PM Form 3.11.2 Informed Consent to Participate in Telemedicine Services
Reference ADHS/DBHS Policy 107