Dear Ms. McKelvey:
The Division of Health Improvement/Quality Management Bureau has completed a compliance survey of the services identified above. The purpose of the survey was to determine compliance with federal and state standards; to assure the health, safety, and welfare of individuals receiving services through the Developmental Disabilities Waiver; and to identify opportunities for improvement. This Report of Findings will be shared with the Developmental Disabilities Supports Division for their use in determining your current and future provider agreements. Upon receipt of this letter and Report of Findings your agency must immediately correct all deficiencies which place Individuals served at risk of harm.

**Determination of Compliance:**
The Division of Health Improvement, Quality Management Bureau has determined your agency is in:

**Compliance with all Conditions of Participation.**
This determination is based on your agency's compliance with CMS waiver assurances at the Condition of Participation level. The attached QMB Report of Findings indicates Standard Level deficiencies identified and requires implementation of a Plan of Correction.

**Plan of Correction:**
The attached Report of Findings identifies the Standard Level and/or Condition of Participation deficiencies found during your agency's compliance review. You are required to complete and implement a Plan of Correction. Your agency has a total of 45 business days (10 business days to submit your POC for approval and 35 days to implement your approved Plan of Correction) from the receipt of this letter.
Submission of your Plan of Correction:
Please submit your agency's Plan of Correction in the space on the two right columns of the Report of Findings. (See attachment "A" for additional guidance in completing the Plan of Correction).

Within 10 business days of receipt of this letter your agency Plan of Correction must be submitted to the parties below:

1. Quality Management Bureau, Attention: Plan of Correction Coordinator
   5301 Central Ave. NE Suite 400 Albuquerque, NM 87108

2. Developmental Disabilities Supports Division Regional Office for region of service surveyed

Upon notification from QMB that your Plan of Correction has been approved, you must implement all remedies and corrective actions to come into compliance. If your Plan of Correction is denied, you must resubmit a revised plan as soon as possible for approval, as your POC approval and all remedies must be completed within 45 business days of the receipt of this letter.

Failure to submit your POC within the allotted 10 business days or complete and implement your Plan of Correction within the total 45 business days allowed may result in the imposition of a $200 per day Civil Monetary Penalty until it is received, completed and/or implemented.

Request for Informal Reconsideration of Findings (IRF):
If you disagree with a finding of deficient practice, you have 10 business days upon receipt of this notice to request an IRF. Submit your request for an IRF in writing to:

QMB Deputy Bureau Chief
5301 Central Ave NE Suite #400
Albuquerque, NM 87108
Attention: IRF request

See Attachment "C" for additional guidance in completing the request for Informal Reconsideration of Findings. The request for an IRF will not delay the implementation of your Plan of Correction which must be completed within 45 total business days (10 business days to submit your POC for approval and 35 days to implement your approved Plan of Correction). Providers may not appeal the nature or interpretation of the standard or regulation, the team composition or sampling methodology. If the IRF approves the modification or removal of a finding, you will be advised of any changes.

Please call the Plan of Correction Coordinator at 505-222-8647 if you have questions about the Report of Findings or Plan of Correction. Thank you for your cooperation and for the work you perform.

Sincerely,

Stephanie R. Martinez de Berenger, M.P.A, GCDF

Stephanie R. Martinez de Berenger, M.P.A, GCDF
Team Lead/Healthcare Surveyor
Division of Health Improvement
Quality Management Bureau
Survey Process Employed:

Entrance Conference Date: October 11, 2011

Present:

**Providence Support Services, Inc**
Jamie Benefield, Owner/Program Director/Service Coordinator
Jody McKeelvey, Executive Director

**DOH/DHI/QMB**
Stephanie R. Martinez de Berenger, M.P.A., GCDF, Team Lead/Healthcare Surveyor
Jennifer Bruns, LBSW, Healthcare Surveyor

Exit Conference Date: October 13, 2011

Present:

**Providence Support Services, Inc.**
Jamie Benefield, Owner/Program Director/Service Coordinator
Jody McKeelvey, Executive Director
Sara Elliott, Quality Assurance Director
John McKeelvey, Administrative Assistant

**DOH/DHI/QMB**
Stephanie R. Martinez de Berenger, M.P.A., GCDF, Team Lead/Healthcare Surveyor
Jennifer Bruns, LBSW, Healthcare Surveyor

Total Homes Visited
Number: 4
- Supported Homes Visited
Number: 4

Administrative Locations Visited
Number: 1

Total Sample Size
Number: 5
- Jackson Class Members
- Non-Jackson Class Members
- Supported Living
- Adult Habilitation
- Community Access

Persons Served Interviewed
Number: 5

Person Served Records Reviewed
Number: 5

Direct Service Professionals Interviewed
Number: 5

Direct Service Professionals Record Review
Number: 28

Service Coordinator Record Review
Number: 1

Administrative Files Reviewed
- Billing Records
- Medical Records
- Incident Management Records
- Personnel Files
- Training Records
- Agency Policy and Procedure
- Caregiver Criminal History Screening Records
- Employee Abuse Registry
• Human Rights Notes and/or Meeting Minutes
• Evacuation Drills
• Quality Assurance / Improvement Plan

CC: Distribution List: DOH - Division of Health Improvement
DOH - Developmental Disabilities Supports Division
DOH - Office of Internal Audit
HSD - Medical Assistance Division
Attachment A

Provider Instructions for Completing the QMB Plan of Correction (POC) Process

Introduction:
After a QMB Compliance Survey, your QMB Report of Findings will be sent to you via e-mail.

Each provider must develop and implement a Plan of Correction (POC) that identifies specific quality assurance and quality improvement activities the agency will implement to correct deficiencies and prevent continued non compliance.

Agencies must submit their Plan of Correction within 10 business days from the date you receive the QMB Report of Findings. (Providers who do not submit a POC within 10 business days will be referred to the Internal Review Committee [IRC] for possible actions or sanctions).

Agencies must fully implement their approved Plan of Correction within 45 business days (10 business days to submit your POC for approval and 35 days to implement your approved Plan of Correction) from the date they receive the QMB Report of Findings. (Providers who fail to complete a POC within the 45 business days allowed shall be referred to the IRC for possible actions or sanctions.)

If you have questions about the Plan of Correction process, call the QMB Plan of Correction Coordinator at 505-222-8647 or email at George.Perrault@state.nm.us. Requests for technical assistance must be requested through your DDSD Regional Office.

The POC process cannot resolve disputes regarding findings. If you wish to dispute a finding on the official Report of Findings, you must file an Informal Reconsideration of Findings (IRF) request within ten (10) business days of receiving your report. Please note that you must still submit a POC for findings that are in question (see Attachment “C”).

Instructions for Completing Agency POC:

Required Content
Your Plan of Correction should provide a step-by-step description of the methods to correct each deficient practice to prevent recurrence and information that ensures the regulation cited is in compliance. The remedies noted in your POC are expected to be added to your Agency’s required, annual Quality Assurance Plan.

If a deficiency has already been corrected, the plan should state how it was corrected, the completion date (date the correction was accomplished), and how possible recurrence of the deficiency will be prevented.

The Plan of Correction must address the required six CMS core elements to address each deficiency of the POC:
1. How the specific and realistic corrective action will be accomplished for individuals found to have been affected by the deficient practice.
2. How the agency will identify other individuals who have the potential to be affected by the same deficient practice, and how the agency will act to protect individuals in similar situations.
3. What QA measures will be put into place or systemic changes made to ensure that the deficient practice will not recur
4. Indicate how the agency plans to monitor its performance to make sure that solutions are sustained. The agency must develop a QA plan for ensuring that correction is achieved and sustained. This QA plan must be implemented, and the corrective action evaluated for its effectiveness. The plan of correction is integrated into the agency quality assurance system; and
5. Include dates when corrective action will be completed. The corrective action completion dates must be acceptable to the State.
6. The POC must be signed and dated by the agency director or other authorized official.

The following details should be considered when developing your POC:
- Details about how and when Consumer, Personnel and Residential files are audited by Agency personnel to ensure they contain required documents;
- Information about how Medication Administration Records are reviewed to verify they contain all required information before they are distributed, as they are being used, and after they are completed;
- Your processes for ensuring that all staff are trained in Core Competencies, Incident Reporting, and Individual-Specific service requirements, etc;
- How accuracy in Billing documentation is assured;
- How health, safety is assured;
- For Case Management Providers, how ISPs are reviewed to verify they meet requirements, how the timeliness of LOC packet submissions and consumer visits are tracked;
- Your process for gathering, analyzing and responding to Quality data; and,
- Details about Quality Targets in various areas, current status, analyses about why targets were not met, and remedies implemented.

*Note: Instruction or in-service of staff alone may not be a sufficient plan of correction.* This is a good first step toward correction, but additional steps should be taken to ensure the deficiency is corrected and will not recur.

**Completion Dates**
- The plan of correction must include a completion date (entered in the far right-hand column) for each finding. Be sure the date is realistic in the amount of time your Agency will need to correct the deficiency; not to exceed 45 total business days.
- Direct care issues should be corrected immediately and monitored appropriately.
- Some deficiencies may require a staged plan to accomplish total correction.
- Deficiencies requiring replacement of equipment, etc., may require more time to accomplish correction but should show reasonable time frames.

**Initial Submission of the Plan of Correction Requirements**
1. The Plan of Correction must be completed on the official QMB Survey Report of Findings/Plan of Correction Form and received by QMB within ten (10) business days from the date you received the report of findings.
2. For questions about the POC process, call the POC Coordinator, George Perrault at 505-222-8647 for assistance.
3. For Technical Assistance (TA) in developing or implementing your POC, contact your local DDSD Regional Office.
4. Submit your POC to George Perrault, POC Coordinator in any of the following ways:
   a. Electronically at George.Perrault@state.nm.us (preferred method)
   b. Fax to 505-222-8661, or
   c. Mail to POC Coordinator, 5301 Central Avenue SW, Suite 400, Albuquerque, NM 87108
5. Do not submit supporting documentation (evidence of compliance) to QMB until after your POC has been approved by the POC Coordinator.
6. QMB will notify you when your POC has been “approve” or “denied.”
   a. During this time, whether your POC is “approved,” or “denied,” you will have a maximum of 45 business days from the date of receipt of your Report of Findings to correct all survey deficiencies.
b. If your POC is denied, it must be revised and resubmitted as soon as possible, as the 45 business day limit is in effect.
c. If your POC is denied a second time your agency may be referred to the Internal Review Committee.
d. You will receive written confirmation when your POC has been approved by QMB and a final deadline for completion of your POC.
7. Failure to submit your POC within 10 business days without prior approval of an extension by QMB will result in a referral to the Internal Review Committee and the possible implementation of monetary penalties and/or sanctions.

**POC Document Submission Requirements**

Once your POC has been approved by the QMB Plan of Correction Coordinator you must submit copies of documents as evidence that all deficiencies have been corrected, as follows.

1. Your internal documents are due within a maximum of 45 business days of receipt of your Report of Findings.
2. You may submit your documents by postal mail (paper hard copy or on a disc), fax, or electronically (scanned and attached to e-mails).
3. All submitted documents must be annotated; please be sure the tag numbers and Identification numbers are indicated on each document submitted. Documents which are not annotated with the Tag number and Identification number may not be accepted.
4. Do not submit original documents; Please provide copies or scanned electronic files for evidence. Originals must be maintained in the agency file(s) per DDSD Standards.
5. In lieu of some documents, you may submit copies of file or home audit forms that clearly indicate cited deficiencies have been corrected, other attestations of correction must be approved by the Plan of Correction Coordinator prior to their submission.
6. For billing deficiencies, you must submit:
   a. Evidence of an internal audit of billing documentation for a sample of individuals and timeframes;
   b. Copies of “void and adjust” forms submitted to correct all over-billed or unjustified units billed identified during your internal audit.

Revisions, Modifications or Extensions to your Plan of Correction (post QMB approval) must be made in writing and submitted to the Plan of Correction Coordinator at QMB, prior to the due date and are approved on a case-by-case basis. No changes may be made to your POC or the timeframes for implementation without written approval of the POC Coordinator.
QMB Determinations of Compliance

- **“Compliance with Conditions of Participation”**
  The QMB determination of “Compliance with Conditions of Participation,” indicates that a provider is in compliance with all ‘Conditions of Participation,’ (CoP) but may have standard level deficiencies (deficiencies which are not at the condition level) out of compliance. The agency has obtained a level of compliance such that there is a minimal potential for harm to individuals’ health and safety. To qualify for a determination of Compliance with Conditions of Participation, the provider must be in compliance with all Conditions of Participation.

- **“Partial-Compliance with Conditions of Participation”**
  The QMB determination of “Partial-Compliance with Conditions of Participation” indicates that a provider is out of compliance with one (1) to three (3) ‘Conditions of Participation.’ This partial-compliance, if not corrected, may result in a serious negative outcome or the potential for more than minimal harm to individuals’ health and safety. The Agency may also have standard level deficiencies (deficiencies which are not at the condition level).

  Providers receiving a repeat determination of ‘Partial-Compliance’ for repeat deficiencies of CoPs may be referred by the Quality Management Bureau to the Internal Review Committee (IRC) for consideration of remedies and possible actions.

- **“Non-Compliant with Conditions of Participation”**
  The QMB determination of “Non-Compliance with Conditions of Participation,” indicates a provider is significantly out of compliance with Conditions of Participation and/or has:
  - Four (4) Conditions of Participation out of compliance.
  - Multiple findings of widespread non-compliance with any standard or regulation with a significant potential for more than minimal harm.
  - Any finding of actual harm or Immediate Jeopardy.

  The Agency may also have standard level deficiencies (deficiencies which are not at the condition level).

  Providers receiving a repeat determination of ‘Non-Compliance’ will be referred by Quality Management Bureau to the Internal Review Committee (IRC) for consideration of remedies and possible actions.
Attachment C

Guidelines for the Provider
Informal Reconsideration of Finding (IRF) Process

Introduction:
Throughout the QMB Survey process, surveyors are openly communicating with providers. Open communication means surveyors have clarified issues and/or requested missing information before completing the review through the use of the signed/dated “Document Request,” or “administrative Needs,” etc. forms. Regardless, there may still be instances where the provider disagrees with a specific finding. Providers may use the following process to informally dispute a finding.

Instructions:
1. The Informal Reconsideration of the Finding (IRF) request must be in writing to the QMB Deputy Bureau Chief within 10 business days of receipt of the final Report of Findings.
2. The written request for an IRF must be completed on the QMB Request for Informal Reconsideration of Finding form available on the QMB website: http://dhi.health.state.nm.us/qmb
3. The written request for an IRF must specify in detail the request for reconsideration and why the finding is inaccurate.
4. The IRF request must include all supporting documentation or evidence.
5. If you have questions about the IRC process, email the IRF Chairperson, Scott Good at scott.good@state.nm.us for assistance.

The following limitations apply to the IRF process:
- The request for an IRF and all supporting evidence must be received within 10 business days.
- Findings based on evidence requested during the survey and not provided may not be subject to reconsideration.
- The supporting documentation must be new evidence not previously reviewed or requested by the survey team.
- Providers must continue to complete their Plan of Correction during the IRF process
- Providers may not request an IRF to challenge the sampling methodology.
- Providers may not request an IRF based on disagreement with the nature of the standard or regulation.
- Providers may not request an IRF to challenge the team composition.
- Providers may not request an IRF to challenge the DHI/QMB determination of compliance or the length of their DDSD provider contract.

A Provider forfeits the right to an IRF if the request is not made within 10 business days of receiving the report and/or does not include all supporting documentation or evidence to show compliance with the standards and regulations.

The IRF Committee will review the request, the Provider will be notified in writing of the ruling; no face-to-face meeting will be conducted.

When a Provider requests that a finding be reconsidered, it does not stop or delay the Plan of Correction process. Providers must continue to complete the Plan of Correction, including the finding in dispute regardless of the IRF status. If a finding is removed or modified, it will be noted and removed or modified from the Report of Findings. It should be noted that in some cases a Plan of Correction may be completed prior to the IRF process being completed. The provider will be notified in writing on the decisions of the IRF committee.
### Standard of Care

**Deficiencies**

**Agency Plan of Correction, On-going QA/QI & Responsible Party**

**Date Due**

<table>
<thead>
<tr>
<th><strong>CMS Assurance – Service Plans: ISP Implementation</strong></th>
<th><strong>Services are delivered in accordance with the service plan, including type, scope, amount, duration and frequency specified in the service plan.</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Tag # 6L14  Residential Case File</strong></td>
<td><strong>Standard Level Deficiency</strong></td>
</tr>
<tr>
<td>Developmental Disabilities (DD) Waiver Service</td>
<td>Based on record review, the Agency failed to maintain a complete and confidential case file in the residence for 1 of 5 Individuals receiving Supported Living Services.</td>
</tr>
<tr>
<td>Standards effective 4/1/2007</td>
<td>The following was not found, incomplete and/or not current:</td>
</tr>
<tr>
<td><strong>CHAPTER 6. VIII. COMMUNITY LIVING SERVICE PROVIDER AGENCY REQUIREMENTS</strong></td>
<td>• Annual ISP (#1)</td>
</tr>
<tr>
<td><strong>A. Residence Case File:</strong> For individuals</td>
<td>Provider: In addition to stating the Plan of Correction for these findings, also please provide your evidence of on-going Quality Assurance / Quality Improvement specific to this tag below this line.</td>
</tr>
<tr>
<td>receiving Supported Living or Family Living, the</td>
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<tr>
<td>Agency shall maintain in the individual’s home a</td>
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<tr>
<td>complete and current confidential case file for</td>
<td></td>
</tr>
<tr>
<td>each individual. For individuals receiving</td>
<td></td>
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<tr>
<td>Independent Living Services, rather than maintaining</td>
<td></td>
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<tr>
<td>this file at the individual’s home, the complete</td>
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<tr>
<td>and current confidential case file for each</td>
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<tr>
<td>individual shall be maintained at the agency’s</td>
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<td>administrative site. Each file shall include the</td>
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<tr>
<td>following:</td>
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<tr>
<td>(1) Complete and current ISP and all supplemental</td>
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<td>plans specific to the individual;</td>
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<td>(2) Complete and current Health Assessment Tool;</td>
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<td>(3) Current emergency contact information, which</td>
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<tr>
<td>includes the individual’s address, telephone</td>
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<tr>
<td>number, names and telephone numbers of residents</td>
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<tr>
<td>Community Living Support providers, relatives, or</td>
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<tr>
<td>guardian or conservator, primary care physician’s</td>
<td></td>
</tr>
<tr>
<td>name(s) and telephone number(s), pharmacy name,</td>
<td></td>
</tr>
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<td></td>
<td></td>
</tr>
</tbody>
</table>
address and telephone number and dentist name, address and telephone number, and health plan;

(4) Up-to-date progress notes, signed and dated by the person making the note for at least the past month (older notes may be transferred to the agency office);

(5) Data collected to document ISP Action Plan implementation

(6) Progress notes written by direct care staff and by nurses regarding individual health status and physical conditions including action taken in response to identified changes in condition for at least the past month;

(7) Physician’s or qualified health care providers written orders;

(8) Progress notes documenting implementation of a physician’s or qualified health care provider’s order(s);

(9) Medication Administration Record (MAR) for the past three (3) months which includes:

(a) The name of the individual;

(b) A transcription of the healthcare practitioners prescription including the brand and generic name of the medication;

(c) Diagnosis for which the medication is prescribed;

(d) Dosage, frequency and method/route of delivery;

(e) Times and dates of delivery;

(f) Initials of person administering or assisting with medication; and

(g) An explanation of any medication irregularity, allergic reaction or adverse effect.

(h) For PRN medication an explanation for the use of the PRN must include:

(i) Observable signs/symptoms or circumstances in which the medication is to be used, and

(ii) Documentation of the effectiveness/result of the PRN delivered.
(i) A MAR is not required for individuals participating in Independent Living Services who self-administer their own medication. However, when medication administration is provided as part of the Independent Living Service a MAR must be maintained at the individual’s home and an updated copy must be placed in the agency file on a weekly basis.

(10) Record of visits to healthcare practitioners including any treatment provided at the visit and a record of all diagnostic testing for the current ISP year; and

(11) Medical History to include: demographic data, current and past medical diagnoses including the cause (if known) of the developmental disability and any psychiatric diagnosis, allergies (food, environmental, medications), status of routine adult health care screenings, immunizations, hospital discharge summaries for past twelve (12) months, past medical history including hospitalizations, surgeries, injuries, family history and current physical exam.
CMS Assurance – Qualified Providers – The State monitors non-licensed/non-certified providers to assure adherence to waiver requirements. The State implements its policies and procedures for verifying that provider training is conducted in accordance with State requirements and the approved waiver.

Tag # 1A20  Direct Support Personnel Training

<table>
<thead>
<tr>
<th>Standard of Care</th>
<th>Deficiencies</th>
<th>Agency Plan of Correction, On-going QA/QI &amp; Responsible Party</th>
<th>Date Due</th>
</tr>
</thead>
<tbody>
<tr>
<td>Developmental Disabilities (DD) Waiver Service Standards effective 4/1/2007</td>
<td>Based on record review, the Agency failed to ensure that Orientation and Training requirements were met for 2 of 28 Direct Service Professionals. Review of Direct Service Professionals training records found no evidence of the following required DOH/DDSD trainings and certification being completed:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CHAPTER 1 IV. GENERAL REQUIREMENTS FOR PROVIDER AGENCY SERVICE PERSONNEL: The objective of this section is to establish personnel standards for DD Medicaid Waiver Provider Agencies for the following services: Community Living Supports, Community Inclusion Services, Respite, Substitute Care and Personal Support Companion Services. These standards apply to all personnel who provide services, whether directly employed or subcontracting with the Provider Agency. Additional personnel requirements and qualifications may be applicable for specific service standards.</td>
<td>• Pre-Service (DSP #59)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C. Orientation and Training Requirements: Orientation and training for direct support staff and his or her supervisors shall comply with the DDSD/DOH Policy Governing the Training Requirements for Direct Support Staff and Internal Service Coordinators Serving Individuals with Developmental Disabilities to include the following: (1) Each new employee shall receive appropriate orientation, including but not limited to, all policies relating to fire prevention, accident prevention, incident management and reporting, and emergency procedures; and (2) Individual-specific training for each individual under his or her direct care, as described in the individual service plan, prior to working alone with the individual.</td>
<td>• Participatory Communication &amp; Choice Making (DSP #47)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Provider: In addition to stating the Plan of Correction for these findings above, also please provide your evidence of on-going Quality Assurance / Quality Improvement specific to this tag below this line.
Department of Health (DOH) Developmental Disabilities Supports Division (DDSD) Policy - Policy Title: Training Requirements for Direct Service Agency Staff Policy - Eff. March 1, 2007 - II. POLICY STATEMENTS:
A. Individuals shall receive services from competent and qualified staff.
B. Staff shall complete individual-specific (formerly known as “Addendum B”) training requirements in accordance with the specifications described in the individual service plan (ISP) of each individual served.
C. Staff shall complete training on DOH-approved incident reporting procedures in accordance with 7 NMAC 1.13.
D. Staff providing direct services shall complete training in universal precautions on an annual basis. The training materials shall meet Occupational Safety and Health Administration (OSHA) requirements.
E. Staff providing direct services shall maintain certification in first aid and CPR. The training materials shall meet OSHA requirements/guidelines.
F. Staff who may be exposed to hazardous chemicals shall complete relevant training in accordance with OSHA requirements.
G. Staff shall be certified in a DDSD-approved behavioral intervention system (e.g., Mandt, CPI) before using physical restraint techniques. Staff members providing direct services shall maintain certification in a DDSD-approved behavioral intervention system if an individual they support has a behavioral crisis plan that includes the use of physical restraint techniques.
H. Staff shall complete and maintain certification in a DDSD-approved medication course in accordance with the DDSD Medication Delivery Policy M-001.
I. Staff providing direct services shall complete safety training within the first thirty (30) days of employment and before working alone with an individual receiving service.
**CMS Assurance – Health and Welfare** – The state, on an ongoing basis, identifies, addresses and seeks to prevent occurrences of abuse, neglect and exploitation. Individuals shall be afforded their basic human rights. The provider supports individuals to access needed healthcare services in a timely manner.

<table>
<thead>
<tr>
<th>Tag # 1A09   Medication Delivery (MAR) - Routine Medication</th>
<th>Standard Level Deficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Developmental Disabilities (DD) Waiver Service Standards effective 4/1/2007</td>
<td>Medication Administration Records (MAR) were reviewed for the months of June, July, August &amp; October 2011.</td>
</tr>
<tr>
<td><strong>CHAPTER 1 II. PROVIDER AGENCY REQUIREMENTS:</strong> The objective of these standards is to establish Provider Agency policy, procedure and reporting requirements for DD Medicaid Waiver program. These requirements apply to all such Provider Agency staff, whether directly employed or subcontracting with the Provider Agency. Additional Provider Agency requirements and personnel qualifications may be applicable for specific service standards.</td>
<td>Based on record review, 2 of 5 individuals had Medication Administration Records, which contained missing medications entries and/or other errors:</td>
</tr>
<tr>
<td><strong>E. Medication Delivery:</strong> Provider Agencies that provide Community Living, Community Inclusion or Private Duty Nursing services shall have written policies and procedures regarding medication(s) delivery and tracking and reporting of medication errors in accordance with DDSD Medication Assessment and Delivery Policy and Procedures, the Board of Nursing Rules and Board of Pharmacy standards and regulations.</td>
<td>Individual #3 August 2011 Medication Administration Records did not contain the diagnosis for which the medication is prescribed:</td>
</tr>
<tr>
<td>(2) When required by the DDSD Medication Assessment and Delivery Policy, Medication Administration Records (MAR) shall be maintained and include:</td>
<td>• Risperdal 1mg (2 times daily)</td>
</tr>
<tr>
<td>(a) The name of the individual, a transcription of the physician’s written or licensed health care provider’s prescription including the brand and generic name of the medication, diagnosis for which the medication is prescribed;</td>
<td>June 2011 Medication Administration Records did not contain the diagnosis for which the medication is prescribed:</td>
</tr>
<tr>
<td>(b) Prescribed dosage, frequency and</td>
<td>• Depakote EC 500mg (1 time daily)</td>
</tr>
<tr>
<td></td>
<td>• Senna Plus (2 times daily)</td>
</tr>
<tr>
<td></td>
<td>Individual #5 October 2011 Medication Administration Records contained missing entries. No documentation found indicating reason for missing entries:</td>
</tr>
<tr>
<td></td>
<td>• Divalprox Sod ER 250mg (3 times daily) – Blank 10/12/2011 (12:00PM)</td>
</tr>
</tbody>
</table>

*Provider:*

In addition to stating the Plan of Correction for these findings above, also please provide your evidence of on-going Quality Assurance / Quality Improvement specific to this tag below this line.

---


Survey Report #: Survey Report #: Q12.02.61285854.METRO.RTN.01
method/route of administration, times and dates of administration;
(c) Initials of the individual administering or assisting with the medication;
(d) Explanation of any medication irregularity;
(e) Documentation of any allergic reaction or adverse medication effect; and
(f) For PRN medication, an explanation for the use of the PRN medication shall include observable signs/symptoms or circumstances in which the medication is to be used, and documentation of effectiveness of PRN medication administered.

(3) The Provider Agency shall also maintain a signature page that designates the full name that corresponds to each initial used to document administered or assisted delivery of each dose;
(4) MARs are not required for individuals participating in Independent Living who self-administer their own medications;
(5) Information from the prescribing pharmacy regarding medications shall be kept in the home and community inclusion service locations and shall include the expected desired outcomes of administrating the medication, signs and symptoms of adverse events and interactions with other medications;

NMAC 16.19.11.8 MINIMUM STANDARDS:
A. MINIMUM STANDARDS FOR THE DISTRIBUTION, STORAGE, HANDLING AND RECORD KEEPING OF DRUGS:

(d) The facility shall have a Medication Administration Record (MAR) documenting medication administered to residents, including over-the-counter medications. This documentation shall include:
   (i) Name of resident;
   (ii) Date given;
   (iii) Drug product name;
(iv) Dosage and form;
(v) Strength of drug;
(vi) Route of administration;
(vii) How often medication is to be taken;
(viii) Time taken and staff initials;
(ix) Dates when the medication is discontinued or changed;
(x) The name and initials of all staff administering medications.

**Model Custodial Procedure Manual**

**D. Administration of Drugs**

Unless otherwise stated by practitioner, patients will not be allowed to administer their own medications. Document the practitioner’s order authorizing the self-administration of medications.

All PRN (As needed) medications shall have complete detail instructions regarding the administering of the medication. This shall include:

- symptoms that indicate the use of the medication,
- exact dosage to be used, and
- the exact amount to be used in a 24 hour period.
<table>
<thead>
<tr>
<th>Tag #</th>
<th>Medication Delivery - PRN Medication</th>
<th>Standard Level Deficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A09.1</td>
<td>Based on record review, the Agency failed to maintain PRN Medication Administration Records which contained all elements required by standard for 3 of 5 Individuals.</td>
<td></td>
</tr>
</tbody>
</table>

**CHAPTER II. PROVIDER AGENCY REQUIREMENTS:** The objective of these standards is to establish Provider Agency policy, procedure and reporting requirements for DD Medicaid Waiver program. These requirements apply to all such Provider Agency staff, whether directly employed or subcontracting with the Provider Agency. Additional Provider Agency requirements and personnel qualifications may be applicable for specific service standards.

**E. Medication Delivery:** Provider Agencies that provide Community Living, Community Inclusion or Private Duty Nursing services shall have written policies and procedures regarding medication(s) delivery and tracking and reporting of medication errors in accordance with DDSD Medication Assessment and Delivery Policy and Procedures, the Board of Nursing Rules and Board of Pharmacy standards and regulations.

(2) When required by the DDSD Medication Assessment and Delivery Policy, Medication Administration Records (MAR) shall be maintained and include:

- a. The name of the individual, a transcription of the physician’s written or licensed health care provider’s prescription including the brand and generic name of the medication, diagnosis for which the medication is prescribed;
- b. Prescribed dosage, frequency and method/route of administration, times and dates of administration;
- c. Initials of the individual administering or assisting with the medication;
- d. Explanation of any medication irregularity;

Based on record review, the Agency failed to maintain PRN Medication Administration Records which contained all elements required by standard for 3 of 5 Individuals.

**Individual #2 August 2011**

Medication Administration Records did not contain the exact amount to be used in a 24 hour period:

- Diastat Acudial 20mg (PRN)
- Ibuprofen 200mg (PRN)
- Acetaminophen 325mg (PRN)

**July 2011**

Medication Administration Records did not contain the exact amount to be used in a 24 hour period:

- Ibuprofen 200mg (PRN)
- Acetaminophen 325mg (PRN)

No Effectiveness was noted on the Medication Administration Record for the following PRN medication:

- Acetaminophen 325mg – PRN – 07/15 (given 2 times)

**June 2011**

Medication Administration Records did not contain the exact amount to be used in a 24 hour period:

- Ibuprofen 200mg (PRN)
- Acetaminophen 325mg (PRN)

No Effectiveness was noted on the Medication Administration Record for the following PRN medication:

Provider: In addition to stating the Plan of Correction for these findings above, also please provide your evidence of on-going Quality Assurance / Quality Improvement specific to this tag below this line.

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Survey Report #: Survey Report #: Q12.02.61285854.METRO.RTN.01
(e) Documentation of any allergic reaction or adverse medication effect; and
(f) For PRN medication, an explanation for the use of the PRN medication shall include observable signs/symptoms or circumstances in which the medication is to be used, and documentation of effectiveness of PRN medication administered.

(3) The Provider Agency shall also maintain a signature page that designates the full name that corresponds to each initial used to document administered or assisted delivery of each dose;

(4) MARs are not required for individuals participating in Independent Living who self-administer their own medications;

(5) Information from the prescribing pharmacy regarding medications shall be kept in the home and community inclusion service locations and shall include the expected desired outcomes of administrating the medication, signs and symptoms of adverse events and interactions with other medications;

NMAC 16.19.11.8 MINIMUM STANDARDS:  
A. MINIMUM STANDARDS FOR THE DISTRIBUTION, STORAGE, HANDLING AND RECORD KEEPING OF DRUGS:  
(d) The facility shall have a Medication Administration Record (MAR) documenting medication administered to residents, including over-the-counter medications. This documentation shall include:
   (i) Name of resident;
   (ii) Date given;
   (iii) Drug product name;
   (iv) Dosage and form;
   (v) Strength of drug;
   (vi) Route of administration;
   (vii) How often medication is to be taken;

<table>
<thead>
<tr>
<th>Individual #3</th>
<th>July 2011</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medication Administration Records did not contain the circumstance for which the medication is to be used:</td>
<td></td>
</tr>
<tr>
<td>• Lorazepam 0.5 mg (PRN)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Individual #4</th>
<th>August 2011</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Effectiveness was noted on the Medication Administration Record for the following PRN medication:</td>
<td></td>
</tr>
<tr>
<td>• Risperidone 0.5mg – PRN – 08/25 (given 1 time)</td>
<td></td>
</tr>
</tbody>
</table>

- Ibuprofen 200mg – PRN – 06/27 (given 1 time)
(viii) Time taken and staff initials;
(ix) Dates when the medication is discontinued or changed;
(x) The name and initials of all staff administering medications.

Model Custodial Procedure Manual
_D. Administration of Drugs_

Unless otherwise stated by practitioner, patients will not be allowed to administer their own medications. Document the practitioner’s order authorizing the self-administration of medications.

All PRN (As needed) medications shall have complete detail instructions regarding the administering of the medication. This shall include:

- symptoms that indicate the use of the medication,
- exact dosage to be used, and
- the exact amount to be used in a 24 hour period.

Department of Health
Developmental Disabilities Supports Division (DDSD) Medication Assessment and Delivery Policy - Eff. November 1, 2006

_F. PRN Medication_

3. Prior to self-administration, self-administration with physical assist or assisting with delivery of PRN medications, the direct support staff must contact the agency nurse to describe observed symptoms and thus assure that the PRN medication is being used according to instructions given by the ordering PCP. In cases of fever, respiratory distress (including coughing), severe pain, vomiting, diarrhea, change in responsiveness/level of consciousness, the nurse must strongly consider the need to conduct a face-to-face assessment to assure that the PRN does not mask a condition better treated by seeking medical attention. This does not apply to home
based/family living settings where the provider is related by affinity or by consanguinity to the individual.

4. The agency nurse shall review the utilization of PRN medications routinely. Frequent or escalating use of PRN medications must be reported to the PCP and discussed by the Interdisciplinary for changes to the overall support plan (see Section H of this policy).

### H. Agency Nurse Monitoring

1. Regardless of the level of assistance with medication delivery that is required by the individual or the route through which the medication is delivered, the agency nurses must monitor the individual’s response to the effects of their routine and PRN medications. The frequency and type of monitoring must be based on the nurse’s assessment of the individual and consideration of the individual’s diagnoses, health status, stability, utilization of PRN medications and level of support required by the individual’s condition and the skill level and needs of the direct care staff. Nursing monitoring should be based on prudent nursing practice and should support the safety and independence of the individual in the community setting. The health care plan shall reflect the planned monitoring of the individual’s response to medication.

**Department of Health Developmental Disabilities Supports Division (DDSD) - Procedure Title:**
**Medication Assessment and Delivery**
**Procedure Eff Date:** November 1, 2006

C. 3. Prior to delivery of the PRN, direct support staff must contact the agency nurse to describe observed symptoms and thus assure that the PRN is being used according to instructions given by the ordering PCP. In cases of fever, respiratory distress (including coughing), severe pain, vomiting, diarrhea,
change in responsiveness/level of consciousness, the nurse must strongly consider the need to conduct a face-to-face assessment to assure that the PRN does not mask a condition better treated by seeking medical attention. (References: Psychotropic Medication Use Policy, Section D, page 5 Use of PRN Psychotropic Medications; and, Human Rights Committee Requirements Policy, Section B, page 4 Interventions Requiring Review and Approval – Use of PRN Medications).

a. Document conversation with nurse including all reported signs and symptoms, advice given and action taken by staff.

4. Document on the MAR each time a PRN medication is used and describe its effect on the individual (e.g., temperature down, vomiting lessened, anxiety increased, the condition is the same, improved, or worsened, etc.).
<table>
<thead>
<tr>
<th>Standard of Care</th>
<th>Deficiencies</th>
<th>Agency Plan of Correction, On-going QA/QI &amp; Responsible Party</th>
<th>Date Due</th>
</tr>
</thead>
</table>

**CMS Assurance – Financial Accountability** – *State financial oversight exists to assure that claims are coded and paid for in accordance with the reimbursement methodology specified in the approved waiver.*

<table>
<thead>
<tr>
<th>Tag # 5I44 Adult Habilitation Reimbursement</th>
<th>Standard Level Deficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Developmental Disabilities (DD) Waiver Service Standards effective 4/1/2007</td>
<td>Based on record review, the Agency failed to provide written or electronic documentation as evidence for each unit billed for Adult Habilitation Services for 2 of 4 individuals.</td>
</tr>
</tbody>
</table>

**MAD-MR: 03-59 Eff 1/1/2004 8.314.1 BI RECORD KEEPING AND DOCUMENTATION REQUIREMENTS:** Providers must maintain all records necessary to fully disclose the extent of the services provided to the Medicaid recipient. Services that have been billed to Medicaid, but are not

**Individual #1**
July 2011
- The Agency billed 120 units of Adult Habilitation (T2021) from 07/20/2011 through 07/26/2011. Documentation received accounted for 96 units.

**Individual #4**
August 2011

Provider:
In addition to stating the Plan of Correction for these findings, also please provide your evidence of on-going Quality Assurance / Quality Improvement specific to this tag below this line.

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Survey Report #: Survey Report #: Q12.02.61285854.METRO.RTN.01
substantiated in a treatment plan and/or patient records for the recipient are subject to recoupment.


CHAPTER 5 XVI. REIMBURSEMENT

A. Billable Unit. A billable unit for Adult Habilitation Services is in 15-minute increments hour. The rate is based on the individual’s level of care.

B. Billable Activities
(1) The Community Inclusion Provider Agency can bill for those activities listed and described on the ISP and within the Scope of Service. Partial units are allowable. Billable units are face-to-face, except that Adult Habilitation services may be non-face-to-face under the following conditions: (a) Time that is non-face-to-face is documented separately and clearly identified as to the nature of the activity; and (b) Non-face-to-face hours do not exceed 5% of the monthly billable hours.

(2) Adult Habilitation Services can be provided with any other services, insofar as the services are not reported for the same hours on the same day, except that Therapy Services and Case Management may be provided and billed for the same hours.
Date:               February 28, 2012
To:                 Ms. Jody McKelvy, Executive Director
Provider:           Providence Support Services, Inc.
Address:            2225 4th Street NW
State/Zip:          Albuquerque, New Mexico   87102
CC:                 Terri McCaslin, Board Chair
                        13505 McCall Court NE
                        Albuquerque, New Mexico   87123
Region:             Metro
Survey Date:        October 11 - 13, 2011
Program Surveyed:   Developmental Disabilities Waiver
Services Surveyed:  Community Living (Supported Living) & Community Inclusion (Adult
                    Habilitation & Community Access)
Survey Type:        Routine

Dear Ms. McKelvy:

The Division of Health Improvement Quality Management Bureau received, reviewed and approved the documents you submitted for your Plan of Correction.

**Your Plan of Correction is closed.**

To maintain ongoing compliance with Standards and regulations, continue to use the Quality Improvement/Quality Assurance processes in your Plan of Correction, including:

- Monthly book reviews are completed by the Quality Assurance Director as part of the agency's QA process.
- The agency maintains a training tracking database. Every payroll, which is every other week, the training tracking database is reviewed by the Executive Director to see what trainings need to be scheduled and hands out scheduled trainings with payroll. The QA Director also maintains her own training tracking to quality assure the employees are trained timely and follows up with the Executive and Program Directors with any concerns or issues.
- The healthcare coordinator and the RN review the MARs as they come in each month and notify the pharmacy of missing entries. The agency residential supervisors, nurse and healthcare coordinator all review the MAR's for missing entries on a weekly/monthly basis. The Quality Assurance Director checks the MARs during the Monthly Book Reviews for all required information as well. The Consultant Pharmacist comes quarterly and reviews MARs.
Consistent implementation of your QA/QI processes will enable you to identify and promptly respond to problems, enhance your service delivery, and result in fewer Deficiencies in future QMB surveys.

Thank you for your cooperation with the Plan of Correction process, and for the work you and your team perform.

Sincerely,

George Perrault, MBA
Plan of Correction Coordinator

Cc: DHI
DDSD