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A.G.Abrashkevich*, S.I.Vinitsky, V.A.Rostovtsev

## SOLVING PROBLEMS

# ON HYDROGEN ATOM 

IN ELECTRIC FIELD
BY MEANS OF REDUCE

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Uzhgorod State University, Uzhgorod, USSR

1. In recent years, atom + field systems have been intensi vely studied both theoretically and experimentally, see, for in stance, review [1]. For a theoretical description it is first necessary to construct a perturbative theory (PT). The calculation by PT is conveniently made by the REDUCE system described in [2].

In our previous papers ( $[3,4]$ ) we made use of the systen for calculating the energies and wave functions of a hydrogen atom in an inhomogeneous electric field. In this note (as well as in preliminary note [9]), we shall consider a homogenious field for simplicity. Our purpose is to demonstrate the possibilities of the REDUCE for solving PT problems in quantum mechanics in the language of the dynamic-symmetry group. Recall that for the hydrogen atom with no field it is the $S O(4)$ group introduced by Fok ([5]).
2. Our approach ([6]) is based on the dynamical group SO(4,2) (see [7]) which takes the external field into account as well. A unitary irreducible representation of the algebra SO(4) is connected by a dilatation transformation with wave functions of the discrete spectrum of a hydrogen atom, and perturbations of a polynomial form are directly expressed through generators of the algebra $S O(4,2)$. Therefore, the solution of an initial spectral problem is reduced to a pure algebraic procedure well adjusted for implementation with the help of REDUCE. Unlike standard PT, here corrections to eigenfunctions in each finite order are expressed by a linear combination of a finite number of the basis functions of a unitary irreducible representation of SO(4,2). Corrections to the eigenvalues and coefficients of that combination are given by polynomials of eigenvalues of a complete set of commuting generators of the SO(4,2) algebra characterizing the unperturbed problem. For the Stark effect such a set [n $\Delta \mathrm{m}$ ] is def ined by parabolic quantum numbers $\left[n_{1} n_{2} m\right]: n=n_{1}+n_{2}+m+1$ is a principal quantum number: $-\Delta=n_{2}-n_{1}$ is a third projection of the Runge-Lenz vector; and $m$ is a third projection of the electron orbital moment on field $\vec{F}$
3. This note presents a short account of an algebraic PT and the text of the program "STARK". This_program implements the PT in the source language of the REDUCE system. It allows us to ob -
tain, in $k$-th order of PT, analytic expressions in the form of polynomials of three variables $\mathrm{n}, \Delta$, and m for corrections to the energy and for coefficients $\mathrm{b}_{\mathrm{st}}^{(\mathrm{k})},-2 \mathrm{k} \leq \mathrm{s}, \mathrm{t} \leq 2 \mathrm{k}$ of the linear combination of basis functions corresponding to the k -th correc tion to the wave function

$$
\psi_{n_{1} n_{2} m}^{(k)}=\sum_{s=-2 k}^{2 k} \sum_{t=-2 k}^{2 k} b_{s t}^{(k)}|s t\rangle
$$

where $|s t\rangle=\left|n_{1}+s, n_{2}+t, m\right\rangle$ is a state vector in $k$-th order of PT. These corrections appear when a hydrogen atom with charge $Z_{a}$ being in an arbitrary state $\left|n_{1} n_{2} m\right\rangle$ is placed in a homogenious field $F$. Hereafter we use atomic units. As a test we present the expressions for $E^{(p)}$ and $b_{s}^{(p)}, p=1,2$. The proposed procedure may be generalized to other polynomial fields and other systems, for instance, to a quantum-mechanical oscillator in an external polynomial field. In the latter case it is necessary to use an osci llator representation of the group $S O(4,2)$ or $S O(2,1)([7])$.

When REDUCE is applied to problems like these, the need is arising to define an action of operators $\hat{\mathrm{f}}(\mathrm{r}, \mathrm{z})$ of two indepen dent variables $r, z$ on a state vector $\langle r, z \mid s t\rangle=\operatorname{KET}(s, t)$. Unlike the article [8], where similar operation in two-dimensional GrossNeveu's model is defined on the basis state 10$\rangle$, we define the action of an operator on an arbitrary state $\left|n_{1} n_{2} \mathrm{~m}\right\rangle$. It allows us to realize degenerate PT shemes.
4. Let us introduce the notation and recall basis properties of the group $S O(4,2)$. The Lie algebra $S O(4,2)$ is formed by $15 \mathrm{ge}-$ nerators $L_{\alpha \beta}=-L_{\beta \alpha}$, where $a, \beta=1, \ldots, 6$ :

$$
\begin{equation*}
\left[\mathrm{L}_{\alpha \beta}, \mathrm{L}_{\alpha \gamma}\right]=\mathrm{ig} \mathrm{~g}_{\alpha \alpha} \mathrm{L}_{\beta \gamma}, \quad g_{\alpha \alpha}=(1111-1-1) \tag{1}
\end{equation*}
$$

In the $x$-representation $L_{\alpha \beta}$ are given by the relations (i, j, $k=1,2,3$ )

$$
\begin{align*}
& L_{i j}=x_{i} p_{j}-x_{j} p_{i} \equiv \varepsilon_{i j k} L_{k} \\
& L_{i 4}=1 / 2\left(x_{i} \vec{p}^{2}+2 i p_{i}-2 \vec{x} p_{i}-x_{i}\right) \equiv A_{i} \\
& L_{i 5}=1 / 2\left(x_{i} \vec{p}^{2}+2 i p_{i}-2 \overrightarrow{x p} p_{i}+x_{i}\right)  \tag{2}\\
& L_{46}=1 / 2\left(r \vec{p}^{2}-r\right), \quad L_{56}=1 / 2\left(r \vec{p}^{2}+r\right), \\
& L_{45}=-i\left(1+\frac{\text { tp }}{}\right), \quad L_{i 6}=-r_{i},
\end{align*}
$$

where $p_{k}=-i \partial_{1} \partial_{k}, L_{k}$ and $A_{k}$ are components of the momentum, orbital moment and the Runge-Lenz vector of the electron, respectively, $\vec{x}=\left(x_{1} x_{2} x_{3}\right)$ are its coordinates, $r=\left(x_{1}^{2}+x_{2}^{2}+x_{3}^{2}\right) 12$. The operators (2) act in the Hilbert space of functions with the scalar product

$$
\begin{equation*}
\langle f \mid g\rangle=\int d^{3} x f^{*}\left(\frac{1}{x}\right) r^{-1} g(\vec{k}) \tag{3}
\end{equation*}
$$

with respect to which they are self-conjugate. For the Stark effect. we choose the basis to be formed of eigenfunctions of three commuting operators $L_{56}, L_{34}$ и $L_{12}$ :

$$
\begin{align*}
& L_{56}\left|n_{1} \bar{n}_{2} m\right\rangle=n\left|n_{1} \bar{n}_{2}^{m}\right\rangle \\
& L_{34}\left|n_{1} \bar{n}_{2} m\right\rangle=\left(n_{2}-n_{1}\right)\left|n_{1} \bar{n}_{2} m\right\rangle  \tag{4}\\
& L_{12}\left|n_{1} \bar{n}_{2} m\right\rangle=m\left|n_{1} \bar{n}_{2} m\right\rangle
\end{align*}
$$

Explicitly, the basis functions $\left\langle\bar{x} \mid n_{1} \bar{n}_{2} \mathrm{~m}\right\rangle$ are given in $[61$. They differ from the Coulomb functions of the hydrogen atom by normalization and in that the argument does not depend on the $\sqrt{-2 E^{(0)}}$ - energy of an unperturbed atom, $E^{(0)}=-z_{2} /\left(2 n^{2}\right)$, i.e. those functions form the Sturm basis with a uniform asymptotics. Their normalization is determined by the scalar product (3), however, we make use of nonnormalized functions

$$
\begin{align*}
& \left|n_{1} n_{2}^{m}\right\rangle=c_{n_{1}}^{-1} n_{2}^{m} \\
& \left.c_{n_{1} n_{2}}=\sqrt{2} n_{1} \bar{n}_{2}^{m}\right\rangle  \tag{5}\\
& \left(n_{1}: n_{2}!/\left[\left(n_{1}+m\right)!\left(n_{2}+m\right)!\right]\right\}^{1 / c}
\end{align*}
$$

The action of the operators $L_{46}$ : $L_{56}$ in this basis is defined by the relations

$$
\begin{aligned}
& \left(L_{46}^{-L} L_{35}\right)\left|n_{1} n_{2}^{m}\right\rangle=\left(n_{1}+m\right)\left|n_{1}-1 n_{2} m\right\rangle+\left(n_{1}+1\right)\left|n_{1}+1 n_{2} m\right\rangle \\
& \left(L_{46} L_{35}\right)\left|n_{1} n_{2} m\right\rangle=\left(n_{2}+m\right)\left|n_{1} n_{2}-1 m\right\rangle+\left(n_{2}+1\right)\left|n_{1} n_{2}+1 m\right\rangle
\end{aligned}
$$

This definition allows us to avoid half-integer degrees of poly nomials when we set the action of the operators $L_{46}$ и $L_{35}$ on the basis, to diminish the number of algebraic substitutions, and to shorten the run time considerably.

The energy E of the hydrogen atom in uniform electric field $F$ may be represented by the expansion

$$
\begin{equation*}
E(F)=E^{(0)}+\sum_{k=1} F^{k} E^{(k)} \tag{7}
\end{equation*}
$$

Then the Scrödinger equation is reduced to the following equation for the state vector $|\phi\rangle$ ([8]):

$$
\begin{equation*}
\left\{L_{56}-n-\sum_{k=1} F^{k} v^{(k)}\left(x_{3}, r\right)\right\}|\phi\rangle=0 \tag{8}
\end{equation*}
$$

where

$$
\begin{aligned}
& v^{(1)}\left(x_{3}, r\right)=\left\{E^{(1)} R-r Z\right\} n / \Omega_{a} \\
& v^{(2)}\left(x_{3}, r\right)=\left\{E^{(k)} r\right\} n / R_{a} \\
& x_{3}=\left(L_{35}-L_{34}\right) n \Omega_{a}, \quad r=\left(L_{56^{-L}} L_{46}\right) n / \Omega_{a} .
\end{aligned}
$$

Accordingly, the solution to (8) is looked for in the form

$$
\begin{equation*}
|\phi\rangle=\left|n_{1} \bar{n}_{2} m\right\rangle+\sum_{k=1}\left|\phi^{(k)}\right\rangle F(k) \tag{9}
\end{equation*}
$$

and instead of (8) we get the system of inhomogeneous equations for $E^{(k)}$ and $|\phi\rangle$ :

$$
\left.L(n)\left|n_{1} n_{2} m\right\rangle=\left\{L_{56}-n\right\} \ln _{1} \bar{n}_{2} m\right\rangle=0
$$

$$
L(n)\left|\phi^{(k)}\right\rangle=v^{(k)}\left(x_{3}, r\right)\left|n_{1} \bar{n}_{2} m\right\rangle+\sum_{p=1}^{k-1} v^{(k-p)}\left(x_{3}, r\right)\left|\phi^{(p)}\right\rangle \equiv f^{(k)}
$$

Taking into consideration the relations (4), (6) and the polynomial form of $V^{(k)}$, we expand the right-hand sides of $f^{(k)}$ and corrections $|\phi(\mathrm{k})\rangle$ over the state vector

$$
\begin{equation*}
|s t\rangle=\left|n_{1}+s n_{2}+t m\right\rangle c_{n_{1} n_{2} m^{m}} \tag{11}
\end{equation*}
$$

normalized so that $|00\rangle=\left|n_{1} n_{2} m\right\rangle$ :

$$
\begin{align*}
& f^{(k)}=\sum_{s=-2 k}^{2 k} \sum_{t=-2 k}^{2 k} f_{s t}^{(k)}|s t\rangle=  \tag{12}\\
& v^{(k)}\left(x_{3}, r\right)|00\rangle+\sum_{p=1}^{k-1} v^{(k-p)}\left|\phi^{(p)}\right\rangle,  \tag{13}\\
& \left|\phi^{(k)}\right\rangle=\sum_{s=-2 k}^{2 k} \sum_{t=-2 k}^{2 k} b_{s t}^{(k)}|s t\rangle, \quad b_{\infty}^{(k)}=0, \quad b_{s}^{(k)}=0 .
\end{align*}
$$

With the orthogonality condition of functions $|s t\rangle$ in the sence of (3) and the relation (4): $L(n) \mid s t)=(s+t) \mid s t)$, obtain, instead of (10),

$$
\begin{align*}
& (s+t) b_{s t}^{(k)}|s t\rangle=f_{s t}^{(k)}|s t\rangle  \tag{14}\\
& f_{\infty}^{(k)}=0
\end{align*}
$$

In each order, starting from the first one, we subsequently obtain $E^{(k+1)}$ and $b_{s t}^{(k+1)}$, by solving an algebraic equation for

$$
\begin{align*}
& f_{00}^{(k+1)}\left(E^{(k+1)}, E^{(p)}, b_{s t}^{(p)}, 1 \leq p \leq k\right)=0  \tag{15}\\
& b_{s t}^{(k+1)}=(s+t)^{-1} f_{s t}^{(k+1)}\left(E^{(p)}, 1 \leq p \leq k+1, b_{s t}^{(1)}, 1 \leq 1 \leq k\right) \tag{16}
\end{align*}
$$

The initial conditions for the recurrence procedure (15). (16) are given by

$$
\begin{equation*}
E^{(0)}=-z_{a}^{2} /\left(2 n^{2}\right), \quad b_{s t}^{(0)}=0 \tag{17}
\end{equation*}
$$

Note that the procedure (15)-(17) fixes the eigenvector $|\phi\rangle$ up to the normalization. In the considered variant of PT the free parameter $b_{00}^{(t)}$ was taken zero. When necessary, the obtained vector $|\phi\rangle$ may be easily normalized with the use of the definition (3).

In this work, the procedure (15)-(17) is realized in the source language of the computer algebra system REDUCE as the program STARK. This program is adjuced in Appendix together with modifications necessary for inhomogenious field case. In the programn we use the following notation:

$$
\begin{aligned}
& X 3 \equiv x_{3}, \quad R \equiv r, \quad F \equiv f^{(k)}, \quad E E \equiv E^{(k)} \\
& B \equiv b_{s t}^{(k)}, \quad K E T(s t) \equiv|s t\rangle \\
& N \equiv n, \quad N 1 \equiv n_{1}, \quad N 2 \equiv n_{2}, \quad M \equiv m, \quad D \equiv \Delta .
\end{aligned}
$$

Note that in the program the set of parabolic numbers [N1 N2 M] in the course of calculations is replaced by an equivalent set [NDM].

Appendix

## Program STARK

COMMENT AT FIRST ONE NEEDS TO READ IN THIS PROGRAM FILE;

OPERATOR X3,R,F,B,EE,KET; NONCOM X3,R;
COMMENT CALCULATE THE PERTURBATION IN THE K-TH ORDER; PROCEDURE V(K, X,Y);

IF $K<1$ THEN 0 ELSE IF $K=1$ THEN
$\operatorname{SUB}(X 1=X, Y 1=Y, \operatorname{EE}(1) * Y 1-Y 1 * X 1)$
ELSE SUB(Y1=Y, EE(K)*Y1);

COMMENT CALCULATE THE RIGHT HAND SIDE OF THE ALGEBRAIC EQUATION IN K-TH ORDER;
FOR ALL $K$ LET $F(K)=$
$\mathrm{N} / \mathrm{ZA*}(\mathrm{~V}(\mathrm{~K}, \mathrm{X} 3, \mathrm{R}) * \mathrm{KET}(0,0)$
FOR P1: $=1: K-1$ SUM V(K-P1, X3, R $) *(F O R S:=-2 * P 1: 2 * P 1$ SUM FOR TT: $=-2 * P 1: 2 * P 1$ SUM $B(P 1, S, T T) * K E T(S, T T))$;

COMMENT FORMAL SUBSTITUTIONS;
LET $X 3=X 3(), R=R()$;
FOR ALL $X, Y$ LET $X 3() * K E T(X, Y)=X 3(X, Y), R() * K E T(X, Y)=R(X, Y)$; FOR ALL $X, Y$ LET $R(X, Y)=((N+X+Y) * K E T(X, Y)-$
$1 / 2 *((X+N 1+M) * K E T(X-1, Y)+$
$(X+N L+1) * K E T(X+1, Y)+$
$(\mathrm{Y}+\mathrm{N} \mathrm{Z}+\mathrm{M}) * \mathrm{KET}(\mathrm{X}, \mathrm{Y}-1)+$
$(Y+N 2+1) * K E T(X, Y+1)) * N / Z A ;$
FOR ALL $X, Y$ LET $X 3(X, Y)=(1 / 2 *(-(X+N 1+M) * K E T(X-1, Y)-$
$(\mathrm{X}+\mathrm{N} 1+1) * \mathrm{KET}(\mathrm{X}+1, \mathrm{Y})+$
$(Y+N Z+M D * K E T(X, Y-1)+$
$(Y+N Z+1) * K E T(X, Y+1))-$
$(\mathrm{Y}-\mathrm{X}-\mathrm{D}) * \mathrm{KET}(\mathrm{X}, \mathrm{Y})) * \mathrm{~N} / 2 \mathrm{Z}$;
COMMENT CALCULATE THE ENERGY AND EXPANSION COEFFICIENTS OF THE STATE VECTOR IN K-TH ORDER;
ARRAY C(1)
PROCEDURE FK(K);
BEGIN SCALAR U,U1,UR;
$U:=F(K) ; U:=D E N$ U; $U:=N U M$ U;
COEFF(U, KET( 0,0$), C)$;
U2: $=C(1)$;
$\operatorname{COEFF}(L E, E E(K), C)$;
U2: $=-\mathrm{UR} \mathrm{C}(1)+\mathrm{EE}(\mathrm{K})$;
WRITE EE(K): =UZ;
FOR I: $=-2 * K: 2 * K$ DO
FOR J: $=-2 * K$ : $2 * K$ DO
<<U2: =IF COEFFCU,KET(I,J),C) NEO 0 THEN C(1)/U1
ELSE 0;
IF ( $I+J$ ) NEO 0 THEN WRITE $B(K, I, J):=U(I+J)$
ELSE WRITE B(K,I, J): $=0$ ) $)$
END;
COMMENT TRANSFORM NUMBERS N1, NZ INTO $N, D, M$;
LET $N 1=(N+D-M-1) / 2, ~ N Z=(N-D-M-1) / 2$;
FOR ALL $X, Y, Z$ SUCH THAT $X<=0 \operatorname{LET} B(X, Y, Z)=0$;

## END OF THE PROGRAM FILE；

COMMENT INPUT OF INITIAL DATA－READ THE DATA UP TO（K－1）－TH ORD； IN STARKIN；

COMMENT OUTPUT OF THE RESULTS UP TO K－TH ORDER TO A FILE．SUBSTI－ TUTE AN INTEGER TO PARAMETER K；
LINELENGTH（65）；OFF NAT，ECHO；OUT STARKOUT；
FKCK）；
WRITE＂END＂；SHUT STARKOUT；ON NAT，ECHO；
END OF RUN；
COMMENT＊＊＊＊＊THE CHANGES NEEDED TO FORM PROGRAM＂MLT＂（INHOMO－ GENIOUS FIELD CASE）；
COMMENT ADD THE LINE：
OPERATOR FAC，A；
COMMENT ADD TWO DEFINITIONS：
FOR ALL X LET FACCX $=$ FOR $I:=1: X$ PRODUCT $I$ ；
PROCEDURE P（N．X）；
If $\mathrm{N}<0$ THEN 0 ELSE If $\mathrm{N}=0$ THEN 1 ELSE If $\mathrm{N}=1$ THEN X
ELSE $((2 * N-1) * X * P(N-1, X)-(N-1) * P(N-2, X)) / N$ ；
COMMENT REPLACE V（K，X，Y）PROCEDURE DEFINITION ：
PROCEDURE V（K，X，Y）；
IF $K<=1$ THEN $O$ ELSE $\operatorname{SUB}(X 1=X, Y 1=Y, E E(K) * Y 1+$
$Z B *(-1) * *(K-1) * Y * * K * P(K-1, X 1 / Y 1)) ;$
COMMENT REPLACE THE DEFINITION OF F（K）；
FOR ALL K LET $F(K)=\mathbb{N} / Z A *(V(K, X 3, R) * K E T(0,0)+$
FOR P1：＝1：K－1 SUM V（K－P1，X3，R）＊
（（FOR S：＝－P1：P1 SUM A（P1，S，－S）＊KET（S，－S））＋ FOR S：＝－P1：P1 SUM

FOR TT：＝－P1：P1 SUM B（P1，S，TT）＊（ET（S，TT）））；
COMMENT REPLACE FKCK PROCEDURE DEFINITION ：
PROCEDURE FKCK）；
BEGIN SCALAR U，U1，U2，K1， 2,21 ；
$\mathrm{U}:=\mathrm{F}(\mathrm{K}) ; \mathrm{U}:=\mathrm{DEN} \mathrm{U} ; \mathrm{U}:=\mathrm{NLM} \mathrm{U} ; \mathrm{K} 1:=\mathrm{K}-2$ ；
COEFF（U，KET $(0,0), C)$ ；UR：$=C(1)$ ：
$\operatorname{COEFF}(U 2, \mathrm{EE}(K), \mathrm{C}) ;$ U2：$=-\mathrm{UR}(\mathrm{C}(1)+\mathrm{EE}(K)$ ；
WRITE EECK）：＝LR；
IF K1＞0 THEN
FOR $\mathrm{S}:=-\mathrm{K} 1: \mathrm{K} 1$ DO
＜＜Z：＝S；Z1：＝－S；U1：＝U1 $\because$ DENCU ；U：＝NUM U；
L2：＝IF COEFF（U，KET（ $Z, Z 1$ ），C）$=0$ THEN 0 ELSE C（1）；
U2：$=I F \operatorname{COEFF}(U 2, A(K 1,2, Z 1), C)=0$ THEN 0
ELSE－पe C（1）+ A $(K 1,2,21)$ ；
WRITE $A(K-2, S,-S):=U 3)>$ ；
FOR I：$=-\mathrm{K}: \mathrm{K}$ DO FOR $J:=-K: K$ D

〈〈U2：＝IF COEFF（U，KET（I，J），C）＝ 0 THEN 0 ELSE C（1）／U1；
IF（I +J ）NEQ 0 THEN WRITE B（K，I，J）：$=\mathrm{U} \mathrm{C}(\mathrm{I}+\mathrm{J})$
ELSE WRITE B（K，I，J）：$=0$ ）
END；
COMMENT ADD THE CONDITIONS：
FOR ALL $X, Y, Z$ SUCH THAT $X<=0$ LET $A(X, Y, Z)=0$ ；
FOR ALL $X, Y, Z$ SUCH THAT $X=1$ LET $B(X, Y, Z)=0$ ；
COMAENT CHANGE THE FILE NAMES IN COMMANDS＂IN＂，＂OUT＂\＆＂SHUT＂； COMMENT $* * * * *$ END OF CHANGES $* * * * *$ ；

COMPENT EXAMPLES OF RESULTS AS THEY have wRITTEN TO THE FILE STARKOUT：
$\operatorname{EE}(1):=(3 * D * N) /(2 * Z A) \leqslant$
$B(1,0,2):=(N * * 3 *(D * * 2+2 * D * M-2 * D * N-4 * D+M * * E-2 * M * N-4 * M$
$+N * * 2+4 * N+3)) /(32 * 2 A * * 3)$
$\operatorname{EE}(2):=(N * * 4 *(3 * D * * 2+9 * M * * 2-17 * N * * 2-19)) /(16 * Z A * * 4) s$
$B(2,0,2):=(N * * 6 *(D * * 4+2 * D * * 3 * M+2 * D * * 3 * N+4 * D * * 3$
$+D * * 2 * M * * 己+6 * D * * Z * M * N+12 * D * * 2 * M-3 * D * * 2 * N * * 2-6 * D * * 2 * N$
$-5 * D * * 2+4 * D * M * * Z * N+8 * D * M * * Z+12 * D * M * N+16 * D * M-4 * D * N * * 3$
$-36 * D * N * * 2-92 * D * N-72 * D+4 * 44 * * 2 * N * * 2+22 * M * * 2 * N+24 * M * * 2$
$-8 * M \times N * * 3-60 * M * N * * 2-136 * M \times N-96 * M+4 * N * * 4+38 * N * * 3$
$+124 * N * * 2+162 * N+72)) /(128 * Z A * * 6) \$$

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Абрашкевич А.Г., Виницкий С.И., Ростовцев В.А. Е4-88-404 Решение задач об атоме водорода
в электрическом поле с использованием системы РЕДЬЮС
Обсуждается методика рещения задач теории возмущений квантовой механики на языке группы динамической симметрии квантовой механики с помощью системы компьютерной алгебры РЕДЬЮС. Приведена программа вычисления в аналитическом виде энергии и волновых функций атома водорода в электрическом поле в произвольном порядке теории возмущений.

Работа выполнена в Лаборатории вычислительной техники и автоматизации ОИЯИ.

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## Abrashkevich A.G., Vinitsky S.I., Rostovtsev V.A.

 Solving Problems on Hydrogen AtomE4-88-404 in Electric Field by Means of REDUCE

A technique of solving perturbation theory problems of quantum mechanics in the language of the dynamic symmetry group by means of REDUCE computer algebra system is discussed. The program for analytical calculation of energy and wave functions of hydrogen atom in an electric field in arbitrary order of the perturbation theory is presented.

The investigation has been performed at the Laboratory of Computing Techniques and Automation, JINR.

