

**Wheat *Mds-1* encodes a heat-shock protein and governs susceptibility towards  
the Hessian fly gall midge**

**Supplementary Material**

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**Table S1.** Hessian fly populations tested against *Mds-1*-knockdown seedlings of Newton wheat.

Biotype/Population	Biotype	Reference
Biotype <i>GP</i>	<i>GP</i> (avirulent to all known R genes)	48
Biotype <i>vH9</i>	<i>vH9</i>	49
Biotype <i>vH13</i>	<i>vH13</i>	49
Biotype <i>L</i>	<i>vH3, vH5, vH6, vH7H8</i>	48
	<i>vH3, vH4, vH5, vH6, vH7H8, vH9, vh10, vH11, vH12,</i>	50
Scott-KS-2005	<i>vH16, vH17, vH18, vH22, vH23, vH24, vH26</i>	
Fannin-TX-2007	<i>vH3, vH4, vH5, vH6, vH7H8, vH9, vH11, vH16, vH17,</i> <i>vH18, vH24, vH26, vH31</i>	50
Kay-OK-2007	<i>vH3, vH4, vH5, vH6, vH7H8, vH9, vh10, vH11, vH12,</i> <i>vH16, vH17, vH18, vH22, vH23, vH24, vH25, vH26,</i> <i>vHdic</i>	50

**Supplementary Table S2. Characterization of transgenic T0 plants with the *Mds-1* knockdown and ectopic expression constructs, respectively\*.**

Plant	Tube Date	Soil Date	Bar	GOIa	GOIb	SS	PHF	PPM	Seed set
<b>Transgenic Bobwhite lines with the <i>Mds-1</i> knockdown construct:</b>									
150	3/30/2009	4/28/2009	+	+	+	+	R	NA	yes
152	3/23/2009	4/28/2009	+	+	+	+	R	NA	yes
358	3/30/2009	6/8/2009	+	+	+	+	R	NA	yes
432	4/23/2009	6/16/2009	+	+	+	+	R	NA	yes
439	5/19/2009	6/16/2009	+	+	+	-	S	NA	yes
523	5/19/2009	6/29/2009	+	+	+	+	R	NA	yes
1630	2/8/2010	2/22/2010	+	+	+	+	R	MR	yes
1639	2/8/2010	3/2/2010	+	+	+	+	R	MR	yes
1703	3/1/2010	3/23/2010	+	+	+	+	R	MR	yes
2030	5/17/2010	6/1/2010	+	+	+	+	R	MR	yes
2059	5/17/2010	6/1/2010	+	+	+	-	S	S	dead
2089	5/24/2010	6/8/2010	+	+	+	+	R	MR	heat dead
2090	5/24/2010	6/8/2010	+	+	+	+	R	MR	yes
2095	5/24/2010	6/8/2010	+	+	+	+	R	MR	yes
2101	5/10/2010	6/8/2010	+	+	+	+	R	MR	heat dead
2129	5/17/2010	6/16/2010	+	+	+	+	R	MR	yes
2164	6/1/2010	6/22/2010	+	+	+	+	R	MR	yes
2222	5/24/2010	6/29/2010	+	+	+	-	S	S	dead
2270	6/1/2010	7/7/2010	+	+	+	-	S	S	yes
2297	7/28/2010	9/8/2010	+	+	+	+	R	MR	yes
2348	8/4/2010	9/15/2010	+	+	+	+	R	MR	yes
2349	8/11/2010	9/22/2010	+	+	+	+	R	MR	yes
2355	6/9/2010	7/19/2010	+	+	+	+	R	MR	yes
2357	6/9/2010	7/19/2010	+	+	+	+	R	MR	yes
<b>Transgenic Molly lines with the <i>Mds-1</i> over-expression construct:</b>									
640	6/15/2009	6/29/2009	+	NA	+	S		yes	
691	6/22/2009	7/6/2009	+	NA	-	R		Disposed	
793	6/22/2009	7/13/2009	+	NA	-	R		yes	
1111	6/29/2009	8/10/2009	+	NA	+	S		yes	
1117	8/10/2009	8/24/2009	+	NA	+	S		yes	
1166	8/10/2009	8/24/2009	+	NA	+	S		yes	
1238	8/17/2009	8/31/2009	+	NA	+	S		yes	
1417	8/31/2009	9/21/2009	+	NA	+	S		yes	
1490	8/17/2009	8/31/2009	+	NA	+	S		yes	

\*Tube date and soil date are the dates when plants were transferred to tubes with culturing media and to pot soil, respectively.

Bar, GOIa, and GOIb represent PCR detection results for the presence of the bar gene (herbicide resistance), inserted gene fragment at antisense, and sense orientation, respectively, in transgenic plants. Positive symbols (+) represent the detection of a DNA fragment with the expected size, whereas negative symbols (-) indicate no PCR amplification at all (Supplementary Figs. S3 and S9b). NA means not applicable or not determined. SS represents successful suppression of *Mds-1* induction by Hessian fly infestation based on northern blot analyses. PHF represents resistant (R) or susceptible (S) phenotypes of the transgenic plants to Hessian fly infestation. Hessian fly biotype GP was used for testing. PPM represents resistant or susceptible phenotypes of transgenic plants to powdery mildew infection. Seeds set represents transgenic plants that produced seeds successfully (yes) or did not reach the seed production stage (dead). MR represents moderate resistance (between grade 1 and 2 of disease severity on a scale of 0 to 4) (51, also see Methods). ‘Heat dead’ represents dead plants due to a malfunction of an overheating growth chamber, whereas ‘disposed’ means plants were disposed before they reached the flowering stage.

**Table S3. Primers used in various experiments.**

Gene Name	Purpose or target	Primer
<b>Primers for <i>Mds-1</i> cloning</b>		
Adapter primer	cDNA cloning	Forward: 5'-AAGCAGTGGTATCAACGCAGAGT
<i>Mds-1</i> R	5' RACE	Reverse: 5'-CGCAAGAACTCGCACACACATC
<i>Mds-1</i> La	cDNA cloning	Forward: 5'-CGTGAAGAAGGAGGAGGTCAAG
Adapter primer	3' RACE	Reverse: 5'-AAGCAGTGGTATCAACGCAGAGT
<i>Mds-1</i> Lb	<i>Mds-1</i> full length	Forward: 5'-ACCTGCGACCCAATCCAGAAC
<i>Mds-1</i> R	Cloning	Reverse: 5'-CGTGAAGAAGGAGGAGGTCAAG
<b>Primers for construct</b>		
<i>Mds-1a'</i>	<i>Mds-1</i> RNAi	Forward: 5'-cacCGTGAAGAAGGAGGAGGTCAAG
<i>Mds-1R</i>	construct	Reverse: 5'-CGCAAGAACTCGCACACACATC
<i>Mds-1</i> Lc	<i>Mds-1</i> expression	Forward: 5'-CTAGtgtatGTGATCGTGCAGCGAG
<i>Mds-1</i> R	construct	Reverse: 5'-CGCAAGAACTCGCACACACATC
<b>Primers for qPCR</b>		
<i>Mds-1</i>	JN162442	Forward: 5'-ACAGCATCCTACCTGCGA Reverse: 5'-ATCGTCGGTGTACTTTG
<i>Hb</i>	3BS_10441917	Forward: 5'-CTACAACCAAAGCAACATC Reverse: 5'-TCAGCAAGAACAGACACAAC
<i>Hd</i>	3DS_2601138	Forward: 5'-CGAGATCTCCGTCTGAGTGA Reverse: 5'-GGCAAGAACAAAGATACTTCAG
<i>HSP16.9</i>	CD896621	Forward: 5'-GTGATGAAATAAACCAAGA Reverse: 5'-GTACAGACTGATGAACAA
<i>HSP16.9</i>	BQ804592	Forward: 5'-TGAGTATTGGGTAGAG Reverse: 5'-AAGAAACGCAATACAGAG
<i>HSP16.9</i>	EU649679	Forward: 5'-TGAGTATTGGGTAGAG Reverse: 5'-AACAAAGACACAACGTAG
<i>HSP16.9</i>	X64618	Forward: 5'-TGAGTATTGGGTAGAG Reverse: 5'-GCAAGAACACAGACACAA
EST	CA717853	Forward: 5'-CGGTCCGATACAGATTTCC Reverse: 5'-CGAAGGGTCAACACAT
EST	CJ618891	Forward: 5'-AGATTAAGGTTAAGGTTCAATTG Reverse: 5'-CTCAACACGGACCAAAT
<i>HSP17.8</i>	AF350423	Forward: 5'-CGACATCAAGGTGCAGGTG Reverse: 5'-GTACTTGGCGTCCTCCTTCTC
Amino acid transporter	BJ275567	Forward: 5'-TCCAATAAACGAACCTAA Reverse: 5'-TGCAAAGGACTTCTTCATG
Enolase	CK208852	Forward: 5'-GATTAGAAATAAAGGCACCGA Reverse: 5'-GTCCGTCAAGCAAAATGT

Methionine synthase	CN012484	Forward: 5'-TGTGATAGTGCCAGTTGA Reverse: 5'-GAAGGACAGGAGTTCATCT
Lipid transfer protein	AF334185	Forward: 5'-TGCTCTGATATGATCTCCAT Reverse: 5'-ATCCTATATGATAAGCGTACATC
Family II extra cellular lipase 1	BQ171153	Forward: 5'-TGAACCTCTAGCCTTTACC Reverse: 5'-CAATGGACGCATGAACAA
Type III peroxidase	CK200808	Forward: 5'-GAAGCATATACTCCTACTCTTG Reverse: 5'-CATGTGAGAGATCAGTTAGTT
Actin	AF326781	Forward: 5'-AAATCTGGCATCACACTTTCTAC Reverse: 5'-GTCTAACATAATCTGGGTAC

**Table S4. Characterization and comparison of *Mds-1*-silenced transgenic wheat lines\*.**

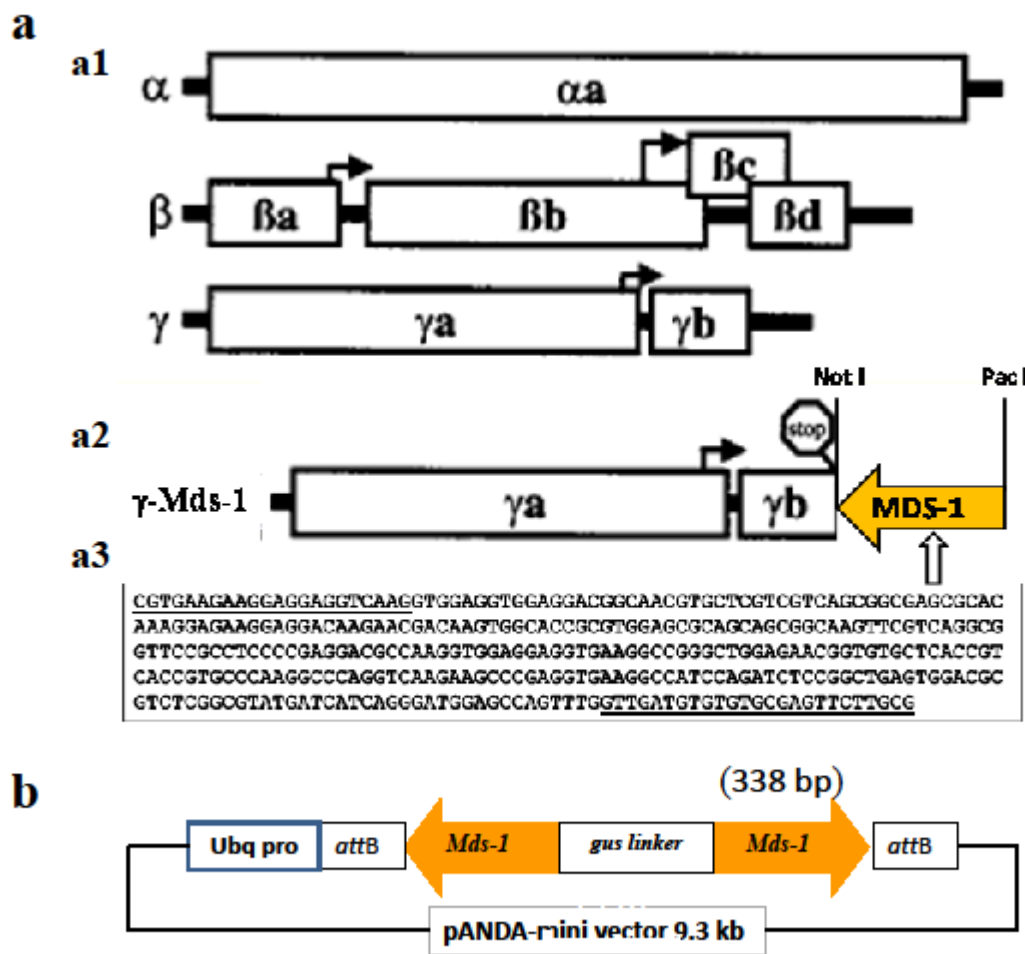
Wheat Lines	Plant Height (cm)		Weight per 100 seeds (g)		Germination Rate (%)		Coleoptile Length (cm)		Radicle Length (cm)	
Bobwhite	75.7	a	4.84	b	93.7	a	8.9	a	14.2	a
T1630	65.3	b	4.46	c	78.7	b	6.7	a	9.2	a
T1639	72	a	4.52	c	84.3	b	6.8	a	12.0	a
T2095	57.7	c	5.54	a	65	c	5.4	b	8.3	c
T2357	74	a	4.16	d	83	b	6.9	a	12.0	a

\*Germination rates were determined by soaking dry seeds in water for 6 hours. Seeds were then transferred to a plastic dish (150x15 mm) with two completely moistened towels at the bottom and two moistened towels on the top of seeds. Dishes were covered to keep moisture and stored under room temperature. Three replicates were carried out with at least 100 seeds for each. Germination data were scored 48 h later. Data were subjected to ANOVA analysis and Tukey's honestly significant difference (HSD) multiple comparison using ProStat Version 5.5(Poly Software International Inc., Pearl River, NY, USA).

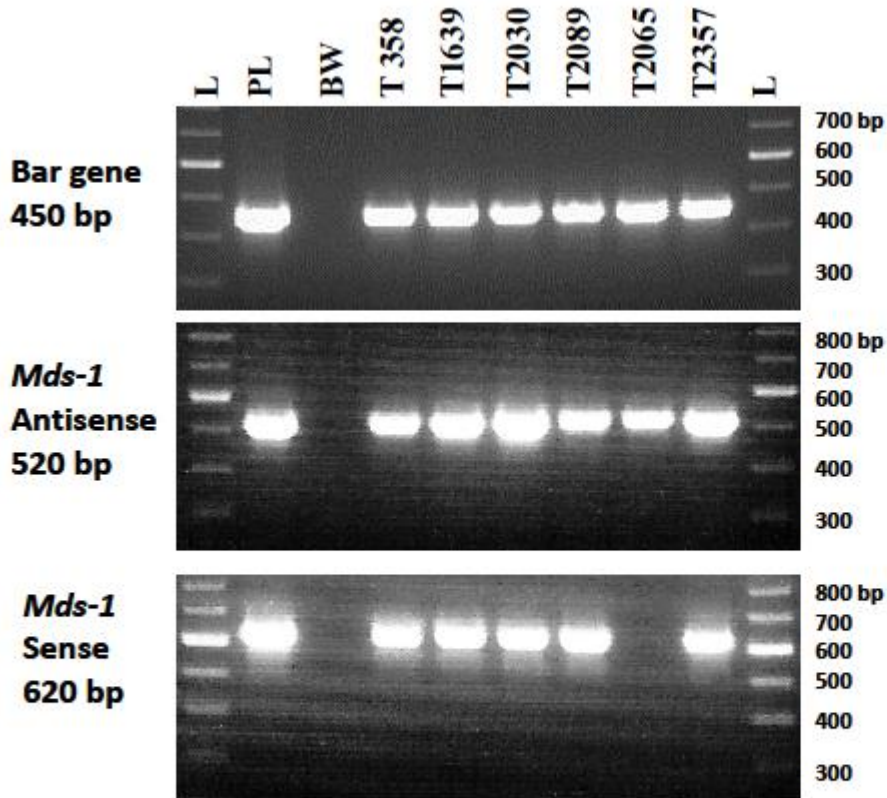
**Figure S1**

AAGCCAGTGTCAACCAAAGTAGACAGCATCTA  
CCTGCGACCCAATCCAGAACAGATTTCGGAGCTAGAAACCAAAGTAACACCGACG  
**ATGTCGATCGTGC**GGAGCAGCGTGGTCAACCCCCTCGCCGACCTCTGGGCTGACCCT  
M S I V R R S S V F D P F A D L W A D P  
  
TTCGACACCTTCCGCTCCATCATCCCAGCGATCTCAGGCAGCAACAGCGAGACGGCCGCG  
F D T F R S I I P A I S G G N S E T A A  
  
TTCGCCAATGCTCGCATGGACTGGAAGGAGACCCCCGAGGCCGACGTCTTCAAGGCCGAC  
F A N A R M D W K E T P E A H V F K A D  
  
CTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGGCAACGTGCTCGTC  
L P G V K K E E V K V E V E D G N V L V  
  
GTCAGCGCGAGCGCACAAAGGAGAACAGGAGGACAAGAACGACAAGTGGCACCGCGTGGAG  
V S G E R T K E K E D K N D K W H R V E  
  
CGCAGCAGCGGCAAGTCGTCAGGCGGTTCCGCCTCCCCGAGGACGCCAAGGTGGAGGAG  
R S S G K F V R R F R L P E D A K V E E  
  
GTGAAGGCCGGCTGGAGAACGGTGTGCTCACCGTCACCGTGCCCAAGGCCAGGTCAAG  
V K A G L E N G V L T V T V P K A Q V K  
  
AAGCCCGAGGTGAAGGCCATCCAGATCTCCGG**TGA**GTGGACGCGTCTGGCGTATGATC  
K P E V K A I Q I S G  
  
ATCAGGGATGGAGCCAGTTGGTTGATGTGTGCGAGTTCTGCGAGTCTGATGAGACA  
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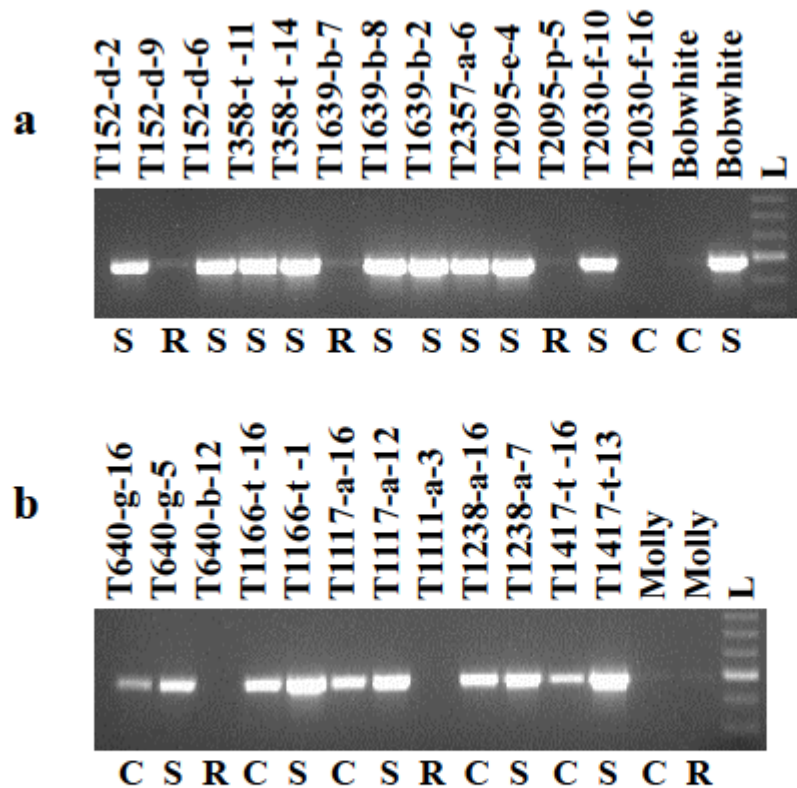
**Figure S2**



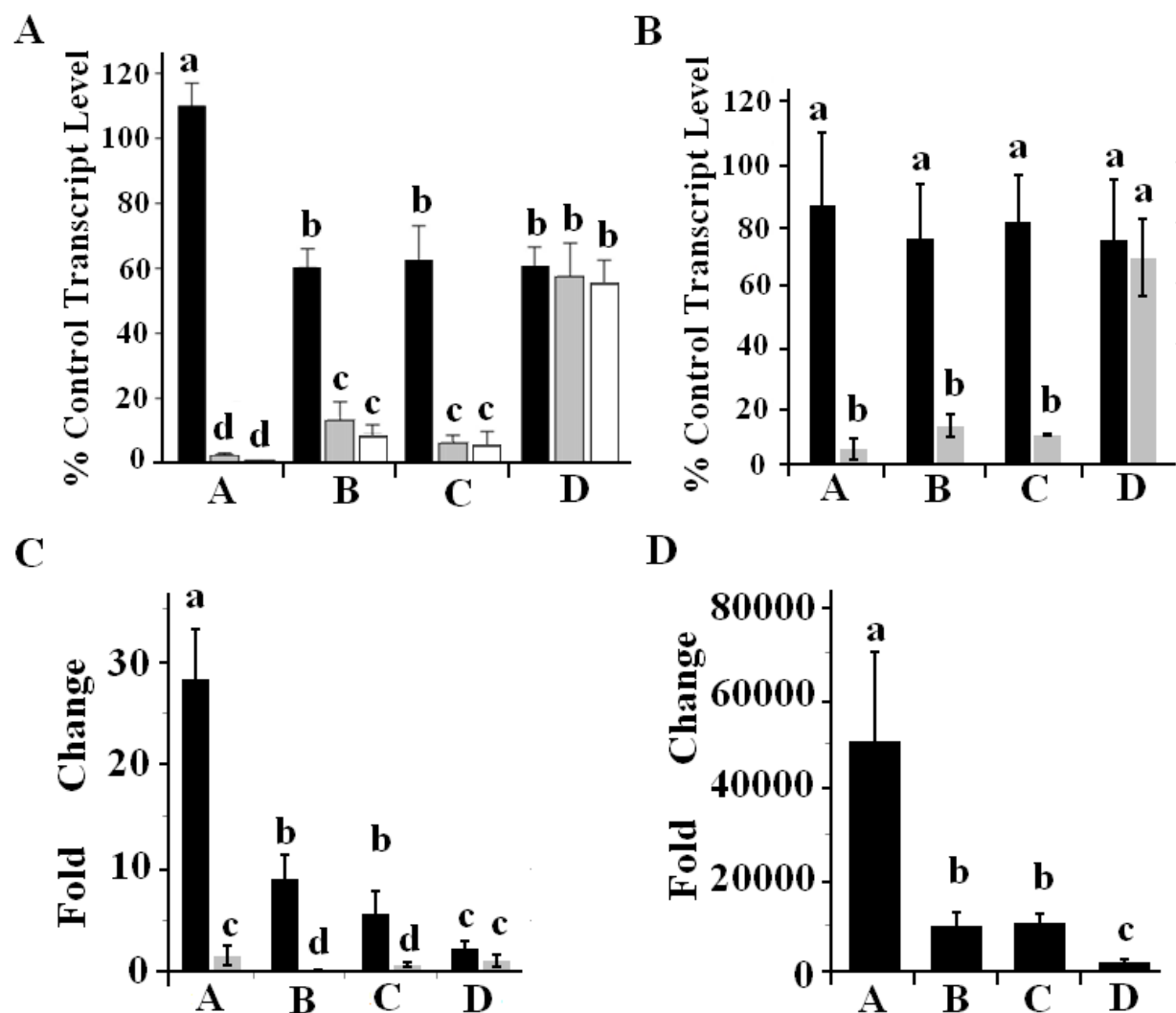
**Figure S3**



**Figure S4**



**Figure S5**



**Figure S6**

**a**

MDS-1	AGCCAGTGTCAACCAAAAGTAG <b>ACAGCATTCTACCTGCGA</b> CCCATCCAGAACAGATTT
CA717853	-----AGCAAAAGCAGACAGCATCTACCTG <b>CGGTCCGATAACAG</b> ----- <b>ATTT</b>
BQ804592	-----GCAACAGCATCTAGCTCTACCTACGATC-----TACAG-----AATT
MDS-1	TTCCCCGAGCTAGAAACCAA----AGTAACACCGACGATGTCGATCGTGC GGAGCAGC
CA717853	<b>TCCC</b> CCGAGCTACAACCAA----AGCAACATCGACAAATGTCGATCGT <b>AGGCGGAGCA</b> <b>AT</b>
BQ804592	TTCCGAGCGAACAGCGAAGCAATCAACACCGACGATGTCGATCGTGC GGAGCAAC
MDS-1	GTGTTCGACCCCTTCGCCGACCTCTGGGC <b>TGACCC</b> TTCGACACCTTCGCTCCAT <b>ATC</b>
CA717853	<b>GTGTTCGACCCCTTCGC</b> CGACCTCTGGGC <b>GACCC</b> CTTCGACACCTTCGCTCCAT <b>CGTC</b>
BQ804592	GTGTTCGACCCCTTCGCCGACCTCTGGGC <b>GACCC</b> CTTCGACACCTTCGCTCCAT <b>CGTC</b>
MDS-1	CCGGCGATCTCAGGCGCAACAGCGAGACGGCCCGTGC <b>CCAA</b> T <b>GCTCG</b> CATGGACTGG
CA717853	CCGGCGATCTCAGGCGCAAC <b>A</b> T <b>GAGAC</b> <b>AGCT</b> CGTGC <b>AA</b> CGCCGGATGGACTGG
BQ804592	CCGGCGATCTC <b>TGGCGCA</b> <b>G</b> AGCGAGACGGCCCGTGC <b>CCAA</b> CGCCGGATGGACTGG
MDS-1	AAGGAGACCCCCGAGGCGCACGTCTTCAAGGCCGACCTCCCCGG <b>CGTGAAGAAGGAGGAG</b>
CA717853	AAGGAGAC <b>CC</b> <b>T</b> GAGGC <b>A</b> ACGTCTTCAAGGCCGACCTCCCCGG <b>GTGAAGAAGGAGGAG</b>
BQ804592	AAGGAGACCCCCGAGGCGCACGTCTTCAAGGCCGACCTCCCCGG <b>GTGAAGAAGGAGGAG</b>
MDS-1	<b>GTCAAGGTGGAGGTGGAGGACGGCAACGTGCTCGTC</b> <b>CAGCGGCGAGCGCACA</b> <b>AAAGG</b> <b>A</b>
CA717853	GTCAAGGTGGAGGTGGAGGACGGCAACGTGCTCGTCAGT <b>GGCGAGCGCACA</b> <b>AAAGG</b> <b>A</b>
BQ804592	GTCAAGGTGGAGGTGGAGGACGGCAACGTGCTCGTCAGCGGCGAGCGCACA <b>AAAGG</b> <b>A</b>
MDS-1	<b>GAAGGAGGACAACAAGACAAGTGGCA</b> ----- <b>CCGC</b> <b>GTGGAGCGCAGCGGGCAAG</b> <b>TT</b> <b>CGTC</b>
CA717853	NAAAGNAACAAGAACGACAAGTGGCAACCGCGTGGAGCGCAAC <b>A</b> <b>AC</b> <b>CG</b> <b>NAAN</b> <b>TTTG</b> <b>TG</b>
BQ804592	GAAGGAGGACAAGAACGACAAGTGGCA-----CCGC <b>GTGGAGCGCAGCGGGCAAG</b> <b>TTTG</b> <b>TGCA</b>
MDS-1	<b>GGCGGTTCCGCTCCCCG</b> ----- <b>AGGACGCCAAGGTGGA</b> ----- <b>GGAGGTGAAGGCCGGGCTGGAGAA</b>
CA717853	GGNG <b>G</b> TTCCGGCTNCCCGAAGAAC <b>C</b> CCAAGGTGGANGAAGGT <b>NAAGGCCGG</b> <b>C</b> <b>T</b> <b>TGAGAA</b>
BQ804592	GGCGGTTCCGGCTGCCGG-----AGGACGCCAAGGTGGA-----GGAGGTGAAGGCCGGGCTGGAGAA
MDS-1	<b>CGGTGTCTCACCGTCACCGTC</b> ----- <b>CCCAAGGCCAGGTCAAGAACGCCCAGGT</b> <b>GAAGGCCA</b>
CA717853	AAGGGTNTCAACGTCAACGTNT <b>CCC</b> <b>AAAGGCCAGGT</b> <b>AAAAAACCCC</b> <b>ANGNAAAGGCCA</b>
BQ804592	CGCG <b>GTGCTCACCGTCACCGTG</b> ----- <b>CCCAAGGCCAGGTCAAGAACGCCCAGGT</b> <b>CAAGGCCA</b>
MDS-1	<b>TCCAGATCTCCGGCTGAGTGGACGCGTCTCGCGTATGATCATCAGGGATGG</b> -----
CA717853	-----CNGATTTCNNNN-----
BQ804592	TCGAGATA <b>TC</b> <b>CCGGCTGAGTGGACGCGTCTCGCGTATGATCATCAGGGATGG</b> -----
MDS-1	----- <b>AGCCAGTTGGTTGATGTTGTCGAGTTCTTGCG</b> ----- <b>AGTCTGATGAGACAT</b> -----
CA717853	-----
BQ804592	TGCAGCCTAAAAGTATGCTACAGT <b>TGTTCTGCGCAGTCTGGT</b> <b>TCACTG</b> <b>TG</b> <b>TGCAAGTTC</b>
MDS-1	----- <b>CTCTGTATTGTGTTCTTCCCCAGT</b> <b>GT</b> <b>TTTCTGTACTTGTGTAATCG</b> <b>GCTAATCG</b> <b>C</b>
CA717853	-----
BQ804592	GGT <b>CTCTGTATTGCGTTCTT</b> <b>TCCCTGTGCGCTGTACTTGTGTAATCG</b> ----- <b>C</b>
MDS-1	CAACAGATT <b>CGCGATGAATAAATGAGAAATAAATTGTTCTGATTTGGGTG</b>
CA717853	-----
BQ804592	CAACAGATT <b>GGCGATGAATAAATGAGAAATAAATTGTTCTGATTTGAGTG</b> -

**b**

*MDS-1*  
AF350423 AAGCCAGTGTCAACCAAAAGTAG**AAGCATCC**TACCTGCCA  
-----CGGGCCGCGACAA-TGGAGGGCAGGATG-----TTGGACTGGAG---

*MDS-1*  
AF350423 T TCCCCGAGCTAGAAC**CAAAC**TAA**CACCGACGAT**GTCG-ATCGTGCGGCGGAGCAG-CG  
- ACCCGCTGATGACGGCGCTGCAGCACCTGCTG-GACGTACCGGACGGCGAGGCCGGCG

*MDS-1*  
AF350423 T GTTCGACCCCTTCGCCGACCTCTGGCTGACCCCTTCGACACCTTCCGCTCCATCATCC  
G ACCCGGQAATGCCGCCGGGAGAAGCAGGCCGACGCCCTACG-TCCG-CGACG

*MDS-1*  
AF350423 C GGCGATCTCAGGGCGGAACAGCGAGACGGCGCGTTCGCCAATGCTCGCATGGACTGGA  
C G-CGGCGCATGGCGGACC-----CCGGCCG-----ACGTGA

*MDS-1*  
AF350423 A GGAGACCCCGAGGCGCACGTC TTCAAGGCCGACCTCCCCGG**CCTCAAGAAGGAGGAGG**  
A GGAGCTGCCGGGCGCTACCGTTCGTGGACATGCCGGGCTGGGGTCCGG**CGACA**

*MDS-1*  
AF350423 T **CAAGGTGGAGGTGGAGGACGGCAAC**CGTCTCGTCAGCGGAGGCCACAAAGGAGA  
T **CAAGGTGAGGTGAGGACGAGCGTGTGGT**GATCAGCGGAGCGGCGGAGGGAGG

*MDS-1*  
AF350423 A **GGAGGACAAGAACGACAAGTGGCACCCGCGTGGAGCGGAGCAGGGCAAGTTCGTCAAGGC**  
A **GAAGG**-----AGCACCCCAAGTACCTCGGATGGAGCGCGATGGCAAGATGATGCGCA

*MDS-1*  
AF350423 G **GTTCCGCCTCCCCGAGGACGCCAAGGTGGAGGGAGGTGAAGGCCGGCTGGAGAACGGTG**  
A GTTCGTGCTCCCCGAGAACGCCGACATGGAGAAGATCTCGGCCGCTGCCGCGACGGCG

*MDS-1*  
AF350423 T **GCTCACCGTCAACCGTGCCAAGGCCAGGTCAAGAACGCCAGGGTAAGGCCATCCAGA**  
T GCTCACCGTCTCCCT-----GGAGAAGCTGCCG-----CCGCCCGAA

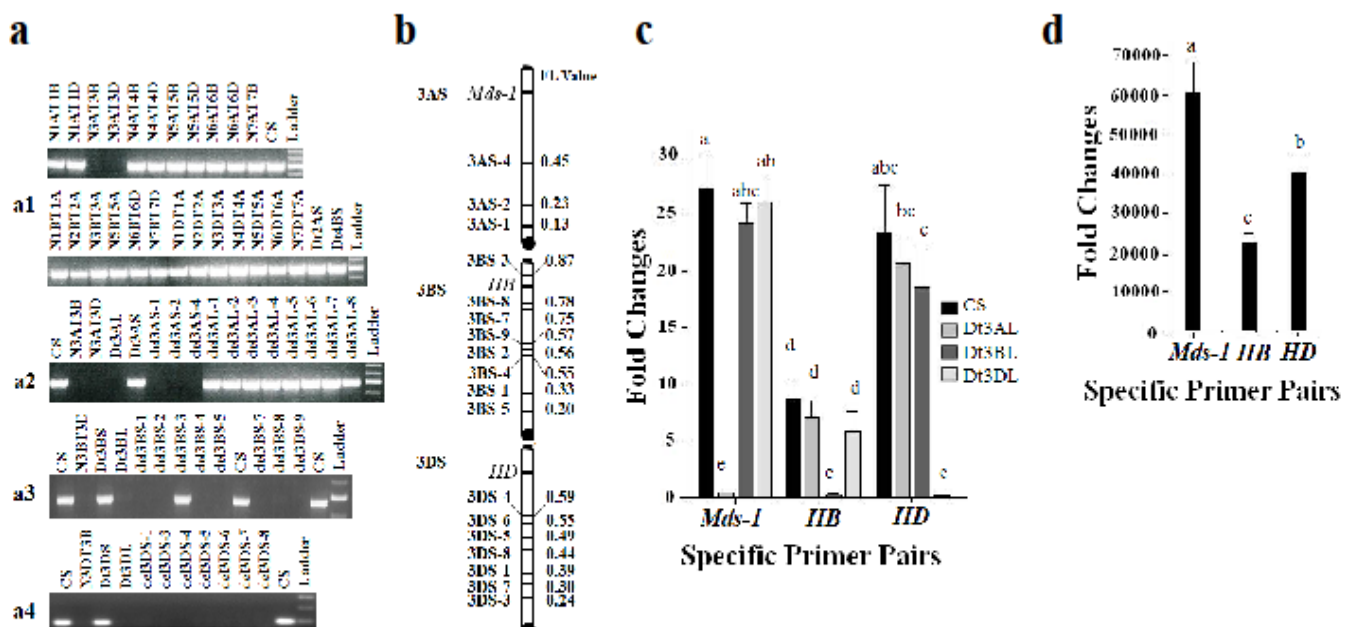
*MDS-1*  
AF350423 T **CTCCGGCTGAGTGGACGCGTCTCGGCGTATGATCATCACGGATGGAGCCAGTTGGT**TG  
A CCAAG-----AAGGCCAAGACCATCCAGGT-----CCAGGTGCGCTG

*MDS-1*  
AF350423 A **TGTGTGTGCGAGTTCTTGCGAGTCTGATGAGACATCTCTGTATTCTGT**TTCTTCCCCA  
A GATGTAT-CGTG-----TGCGCGTCG-----AATCGAAGCAGACAGTAGC-----

*MDS-1*  
AF350423 G TGTTTTCTGTACTTGTGTAATCGGCTAATCGCCAACAG-ATTGGCGATGAATAAATGA  
G AGTTTCCGTG-ATGAGCGATGGTCTGTCTGTCTGGTTTCG-----TTAATGG

*MDS-1*  
AF350423 G AAATAAAATTGTTCTGATTTGGTGC-----AAATGG-----CGATGCCGCCGCG

**Figure S7**



**Figure S8**

**a**

<b>MDS-1</b>	MSIVRRSSVFDPFADLWADPFDTFRS <b>I</b> PAISGGNS---ETAAFANAR <b>MDW</b> KETPEAHVF
<b>HB</b>	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGGNN---ETAAFANAR <b>MDW</b> KETPEAHVF
<b>HD</b>	-----PFADLWADPFDTFRSIVPAISGG <b>S</b> ---ETAAFANAR <b>VDW</b> KETPEAHVF
Contig17	MSIVRRSSVFDPFADLWADPFDTFRS <b>I</b> PAISGGNS---ETAAFANAR <b>MDW</b> KETPEAHVF
Contig4a	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGGNS---ETAAFANAR <b>MDW</b> KETPEAHVF
Contig1	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGGNS---ETAAFANA <b>H</b> DWKETPEAHVF
Contig14r	MSIVRRSNVFNPFAIDLWADPFDTFRS IVPAISGGNS---ETAAFANAR <b>VDW</b> KES <b>P</b> EAHVF
Contig15r	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGGNS---ETAAFANAR <b>VDW</b> KETPEAHVF
Contig11	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGG <b>S</b> ---ETAAFANAR <b>VDW</b> KETPEAHVF
Contig7	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGG <b>SS</b> ---ETAAFA <b>S</b> ARV <b>VDW</b> KETPEAHVF
Contig1ar	MSIVRRSNVFDPFADLWADPFDTFRS IVPAISGG <b>S</b> ---ETAAFANAR <b>VDW</b> KETPEAHVF
Contig10	MSMVRSSNVFDPFADLWADPFDTFRSIVPA <b>MISGN</b> NNDETAAFANAR <b>VDW</b> KETPEAHVF
	*****:***. . * . *****.*:*****:*****
<b>MDS-1</b>	KADLPGVKKEEVKVEVEDGNVLVVSGERTKEKDKN <u>D</u> KWHRVERSSGKFVRRFLPEDAK
<b>HB</b>	KADLPGVKKEEVKVEVEDGNVLVVSGERTKEKD <u>R</u> NDK <u>W</u> HRVERSSGKFVRRFLPEDAK
<b>HD</b>	KADLPGVKKEEVKVEVEDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig17	KADLPGVKKEEVKVEVEDGNVLVVSGERTKEKDKN <u>D</u> KWHRVERSSGKFVRRFLPEDAK
Contig4a	KADLPGVKKEEVKVEVEDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig1	KADLPGVKKEEVKVEVEDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig14r	KADLPGVKKEEVKVE <u>L</u> EDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig15r	KADLPGVKKEEVKVEVEDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig11	KADLPGVKKEEVKVEVEDGNVLVVSGER <u>S</u> RE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig7	KADLPGVKKEEVKVEVEDGNVLVVSGER <u>S</u> RE <u>E</u> DKNDK <u>W</u> HRVERSSGKF <u>A</u> RRFLPEDAK
Contig1ar	KADLPGVKKEEVKVEVEDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
Contig10	KADLPGVKKEEVKVEVEDGNVLVVSGERTKE <u>E</u> DKNDK <u>W</u> HRVERSSGKFVRRFLPEDAK
	*.*****:*****:*****:*****:*****. :*** ***
<b>MDS-1</b>	VEEVKAGLENGVLTVTVPKA <b>Q</b> VKKPEVKAI <b>Q</b> ISG
<b>HB</b>	VEEVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> ISG
<b>HD</b>	V <b>G</b> EVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> IS <b>V</b>
Contig17	VEEVKAGLENGVLTVTVPKA <b>Q</b> VKKPEVKAI <b>Q</b> ISG
Contig4a	VEEVKAGLENGVLTVTVPKAEVKKP <b>Q</b> VKAIEISG
Contig1	VEEVKAGLENGVLTVTVPKAEVKKP <b>Q</b> VKA <b>Q</b> ISG
Contig14r	VEEVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> ISG
Contig15r	V <b>G</b> EVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> IS <b>V</b>
Contig11	VEEVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> ISG
Contig7	VEEVKAGLENGVLTVTVPKAHVKKPEVKAI <b>E</b> ISG
Contig1ar	VEEVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> ISG
Contig10	VEEVKAGLENGVLTVTVPKAEVKKPEVKAI <b>E</b> <b>T</b> <b>G</b>
	* *****:*****:*****:*****:*****. :*** ***

**b**

<b>Mds-1</b>	-----AAGCCAGTGTCAACCAAAG-TAG----- <b>ACAGCATCCTA</b> ----- <b>CCTGCG</b> 40
<b>Hb</b>	AACTCAAATCAGCGTCAAGCAAAG-CAG-----ACAGCATCCT-----AC 40
<b>Hd</b>	-----
Contig17	-----AAGCCAGTGTCAACCAAAG-TAG-----ACAGCATCCTA-----CCTGCG 40
Contig11	-----AAGCCAGTGTCAACGCAACAG-CAA-----ACAACATCCTTT-----CCTAC- 41
Contig14r	-----AAATCAGTGTCAAGCAAAG-TAG-----ACGACATCCTATT-----CCTAC- 41
Contig1	-----AAGCCAGTGTCAAGCAAAG-CAG-----ACAAAATCCCATTCCCATTCCTAC- 49
Contig15r	-----AAGCCAGTGTCAAGCAAAG-CAG-----ACAGCAAGGCCGTT-----CCTAC- 42

Contig4a	-----ATCCCAAACTCAAGCCAGCGTCAA---	GCAAAATCCTACG-----	37
Contig10	-----AAGCCAGTGTCAAGCAAAG-CAAGCAAACAGCATCCTGTT-----	CCTAC	45
Contig1ar	-----ACAGCATCCTAGT-----	CCTAC	18
Contig7	-----AAGCCAGTGTCAACCAAAG-CAG---ACAGCAACCCCAA-----	CCTAC	41

<b>Mds-1</b>	<b>ACCCAATCCAGAACAG</b> ---	ATTTT <u>TTC</u> <u>CC-CG</u> <u>G</u> -- <b>AG</b> CTAG--AAAC <b>CAAAG</b> - <b>TAACACC</b>	90
<b>Hb</b>	CTGCGGTCCGATACAG--	ATTTT <u>CCCC</u> <u>GGAG</u> <u>G</u> -- <b>TACAACCCAAAGCAA</b> -- <b>CATC</b>	89
<b>Hd</b>	-----	-----	
Contig17	ACCCAATCCAGAACAG---	ATTTT <u>TTC</u> <u>CC-CG</u> <u>G</u> -- <b>AG</b> CTAG--AAACCAAAG-TAACACC	90
Contig11	CTACGATCCGATACC <u>G</u> -- <b>AA</b> <u>TTT</u> <u>TTC</u> <u>CGAGCG</u> --	CACAAGCCAAACCAAAGCAAACACT	95
Contig14r	CTACGATCC--ACAG <u>G</u> -- <b>AA</b> <u>TTT</u> <u>TTC</u> <u>CGAGCG</u> --	CACAAGCCGAAGCAAT--CAGCAC	90
Contig1	CTACGATCCAATACAT--	AATCT <u>TTC</u> <u>CC-GAGC</u> -TACAAGCCAAAGCAAG--CAATACC	102
Contig15r	CTACGATCAAATACAT--	AATTTC <u>CCCCCAAG</u> CTACAAGCCAAAGCAAG--CAACACC	96
Contig4a	-TACGATCCGATACAG <u>G</u> -- <b>AA</b> <u>TTT</u> <u>TTC</u> <u>CCCCCG</u> --	AGCTAC--AAACCGAAG-CAACACC	86
Contig10	CTACGATCCGATACAG <u>GAA</u> <u>TTT</u> <u>TTC</u> <u>TTCCGAGCG</u> --	CACAAGCCGAAGCAAT--CAGCAC	101
Contig1ar	CTACGATC--TACAG <u>G</u> -- <b>AA</b> <u>TTT</u> <u>TTC</u> <u>CGAGCG</u> --	CACAAGCCGAAGCAAT--CAACACC	67
Contig7	CTACGATCCAATACAG <u>G</u> -- <b>AA</b> <u>TTT</u> <u>TTC</u> <u>CTAGCG</u> --	CAAGAGTCAAAGCAAG--CAACACC	93

<b>Mds-1</b>	<b>GACGAT</b> TGCGATCGTGC <del>GG</del> GGAGCAGCGT <del>GG</del> TCGACCC <del>CC</del> TTGCCGACCTCTGGGCTGA	150
<b>Hb</b>	GACA <b>ATG</b> TGCGATCGTGAGGCCGAGCAATGTGTT <del>TC</del> GA <del>CC</del> CTT <del>CC</del> GCCGACCTCTGGGCGGA	149
<b>Hd</b>	-----	-----
Contig17	GACG <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	150
Contig11	GACG <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>TT</del> GGGCGGA	155
Contig14r	GACC <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	150
Contig1	GGCA <b>ATG</b> TGCGATCGTGAGGCCGAGTAACGT <del>GG</del> TCAT <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	162
Contig15r	GACA <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	156
Contig4a	GAGA <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	146
Contig10	GACA <b>ATG</b> TGCGATGGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	161
Contig1ar	GACG <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	127
Contig7	GGCA <b>ATG</b> TGCGATCGTGAGGCCGAGCAACGT <del>GG</del> TCAC <del>CC</del> CTT <del>CC</del> GCCGACCT <del>CT</del> GGGCGGA	153

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<b>Mds-1</b>	CCCTTCGACACCTTCCGCTCCATCATCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	202
<b>Hb</b>	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAAAT-----	201
<b>Hd</b>	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	75
Contig17	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	202
Contig11	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	207
Contig14r	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	202
Contig1	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	214
Contig15r	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	208
Contig4a	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGCAACAGC-----	198
Contig10	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGATGATCTCAGGCAACAAACAACAGA	221	
Contig1ar	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCG---GCAGCAG---	178
Contig7	CCCCTCGACACCTTCCGCTCCATCGTCCC <del>GG</del> CGAT--	CTCAGGCCGAGCAGCAGCAG---	207

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<b>Mds-1</b>	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGAC <del>CC</del> CGAGGCCGAC <del>GT</del>	261
<b>Hb</b>	-GAGACAGCT <del>GG</del> T <del>GG</del> CGCAACGCC <del>GG</del> GAT <del>GG</del> ACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	260
<b>Hd</b>	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	134
Contig17	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	261
Contig11	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	266
Contig14r	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	261
Contig1	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	273
Contig15r	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	267
Contig4a	-GAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	257
Contig10	CGAGACAGCT <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	281
Contig1ar	CGAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAA <del>TG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	238
Contig7	CGAGACGGCCGCG <del>GG</del> T <del>GG</del> CGCCAGT <del>GG</del> CTCGCATGGACT <del>GG</del> AGGA <del>GG</del> AGACGCC <del>GT</del> GA <del>GG</del> GCAC <del>GT</del>	267

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**Mds-1** CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 321  
**Hb** CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 320  
**Hd** CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 194  
Contig17 CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 321  
Contig11 CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 326  
Contig14r CTTCAAGGCCGACATCTCCAGGGTGAAGAAGGAGGAGGTCAAGGTGAACTGGAGGACGG 321  
Contig1 CTTCAAGGCCGACCTCCCAGGGTGAAGAAGGAGGAGGTCAAGGTGAACTGGAGGACGG 333  
Contig15r CTTCAAGGCCGACCTCCCAGGGTGAAGAAGGAGGAGGTCAAGGTGAGGTGGAGGACGG 327  
Contig4a CTTCAAGGCCGACCTCCCTGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 317  
Contig10 CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 341  
Contig1ar CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 298  
Contig7 CTTCAAGGCCGACCTCCCCGGCGTGAAGAAGGAGGAGGTCAAGGTGGAGGTGGAGGACGG 327  
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**Mds-1** CAACGTGCTCGTCGTCAAGCAGCGCACAAAGGAGAAGGAGGACAAGAACGACAAGTG 381  
**Hb** CAACGTGCTCGTCGTCAAGCAGCGCACAAAGGAGAAGGAGGACAAGAACGACAAGTG 380  
**Hd** CAACGTGCTCGTCGTCAAGCAGCGCACAAAGGAGGAGGAGGACAAGAACGACAAGTG 254  
Contig17 CAACGTGCTCGTCGTCAAGCAGCGCACAAAGGAGAAGGAGGACAAGAACGACAAGTG 381  
Contig11 CAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGACAAGAACGACAAGTG 386  
Contig14r CAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGACAAGAACGACAAGTG 381  
Contig1 CAAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGACAAGAACGACAAGTG 393  
Contig15r CAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGACAAGAACGACAAGTG 387  
Contig4a CAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGACAAGAACGACAAGTG 377  
Contig10 CAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGAGGACAAGAACGACAAGTG 401  
Contig1ar CAACGTGCTCGTCGTCAAGCAGCGAGCGCACAAAGGAGGACAAGAACGACAAGTG 358  
Contig7 CAACGTGCTCGTCGTCAAGCAGCGAGCGAGGGAGAAGGAGGACAAGAACGACAAGTG 387  
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**Mds-1** GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 441  
**Hb** GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 440  
**Hd** GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 314  
Contig17 GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 441  
Contig11 GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 446  
Contig14r GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 441  
Contig1 CACCCCGCTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGACTTCCGCCTGCCGGAGGACGC 453  
Contig15r GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTGCCGGAGGACGC 447  
Contig4a GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGCTTCCGCCTGCCGGAGGACGC 437  
Contig10 GCACCGCGTGGAGCGTAGCAGCGCAAGTTCGTCAAGCGGCTTCCGCCTGCCGGAGGACGC 461  
Contig1ar GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGTTCCGCCTCCCCGAGGACGC 418  
Contig7 GCACCGCGTGGAGCGCAGCAGCGCAAGTTCGTCAAGCGGCTTCCGCCTGCCGGAGGACGC 447  
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**Mds-1** CAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 501  
**Hb** CAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 500  
**Hd** CAAGGTGGGGGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 374  
Contig17 CAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 501  
Contig11 CAAGGTGGAGGAGGTCAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 506  
Contig14r CAAGGTGGAGGAGGTCAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 501  
Contig1 CAAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 513  
Contig15r CAAGGTGGGGGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 507  
Contig4a CAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 497  
Contig10 CAAGGTGGAGGAGGTGAAGGCCGGTCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 521  
Contig1ar CAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 478  
Contig7 CAAGGTGGAGGAGGTGAAGGCCGGGCTGGAGAAGCGGTGTGCTCACCGTCACCGTGCCCAA 507  
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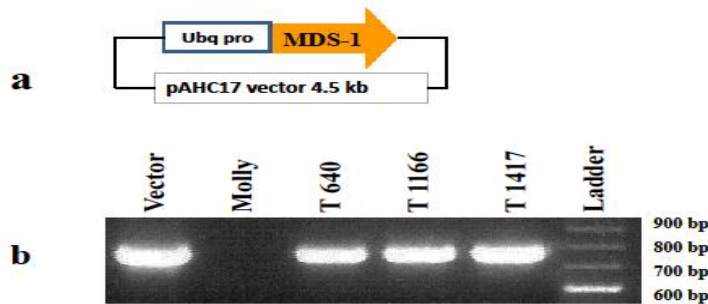
▼  
**Mds-1** GGCCCAGGTCAAGAAGCCCAGGTGAAGGCCATCCAGATCTCCGGCT**TGAG**TGGACCGTC 561  
**Hb** GGCGAGGTCAAGAAGCCCAGGTGAAGGCCATCGAGATCTCCGGCT**TGAG**----- 550  
**Hd** GGCGAGGTCAAGAAGCCCAGGTGAAGGCCAT**CGAGATCTCCGTCTGAGTGA**ATCGTC 434  
Contig17 GGCCCAGGTCAAGAAGCCCAGGTGAAGGCCATCCAGATCTCCGGCT**TGAG**TGGACCGTC 561  
Contig11 GGCGAGGTCAAGAAGCCCAGGTGAAGGCCATCGAGATCTCCGGCT**TGAG**TGCATCGTT 566  
Contig14r GGCGAGGTCAAGAAGCCCAGGTGAAGGCCATCGAGATCTCCGGCT**TGAG**TGCATCGTT 561

Contig1	GGCAGAGGTCAAGAAGCCCCAGGTGAAGGCCATCCAGATCTCCGGCT <b>TGAGTGGACCGTC</b>	573
Contig15r	GGCCGAGGTCAAGAAGCCCCAGGTGAAGGCCATCGAGATCTCCGCT <b>TGAGTGAATGCGTC</b>	567
Contig4a	GGCAGAGGTCAAGAAGCCCCAGGTGAAGGCCATCGAAATCTCCGGCT <b>TGAGTGGACCGTA</b>	557
Contig10	GGCCGAGGTGAAGAAGCCTGAGGTGAAGGCCATCGAGATCACCGGC <b>TGAG-----</b>	571
Contiglar	GGCCGAGGTCAAGAAGCCCCAGGTGAAGGCCATCGAGATCTCCGGCT <b>TGAGTGGACCGTC</b>	538
Contig7	GGCCCACGTCAAGAAGCCCCAGGTGAAGGCCATCGAGATATCCGGCT <b>TGAG-----</b>	557
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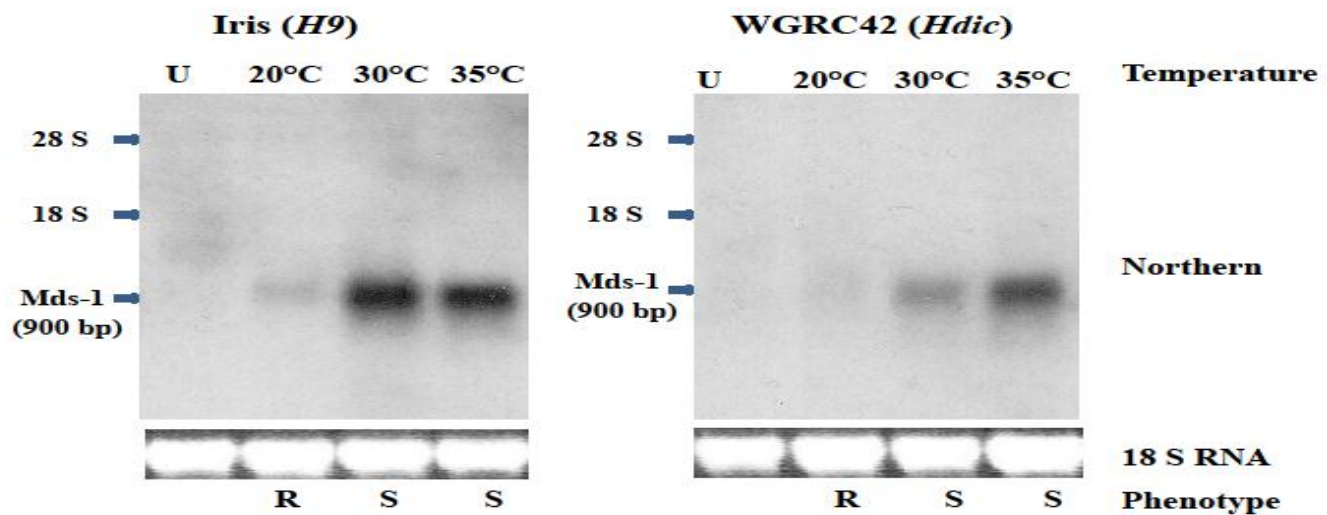
<b>Mds-1</b>	TCGGCG-----TATGATCATCAGGGATGGAGCCAGTTGGTTG <b>ATGTGTGTGCGAG</b> ---	612
<b>Hb</b>	-----TATTGGGTCAAGAGCCTGCTGGTTCATGTGTGCGAGCTC	592
<b>Hd</b>	TCTGCG-----AATGATCATCGGGGTTGGAGCCAGTTGTTATGCAAC <b>CTGAAAG</b> ---	485
Contig17	TCGGCG-----TATGATCATCAGGGATGGAGCCAGTTGGTTATGTTGAG---	612
Contig11	TGGACG-----TATGCTCATTCAAGGTTGGAGCCAATTGGTTGATGTGTGCGAG---	617
Contig14r	TGGACG-----TATGCTCATTCAAGGTTGGAGCCAATTGGTTGATGTGTGCGAG---	612
Contig1	TCGGCG-----TATGATCATCAGGGATGGAGCCAGTTGGTTGATGTGTGCGAGGTC	627
Contig15r	TCTGCG-----AATGATCATCGGGGTTGGAGCCAGTTGTTATGCAACCTGAAAG---	618
Contig4a	TCGTCACGTATGATGATCTTGGGCCGAGCCAGTTGGTTGATGTGTGTAAGAGTTC	617
Contig10	-----CATTGG-----AGCCAGTTAGTTGATGTGTGTTGCGAGGTC	607
Contiglar	TCGGCG-----TATGATCATCAGGGATGGAGCCAGTTGGTTGATGTGTGCGAGGTC	592
Contig7	-----CAT-----TGGAGCCAGTTGATTGACGTGTGCGAGGTC	593
	:* *****:.* * .** : .. *...**	

<b>Mds-1</b>	----- <b>TTCTTG</b> ----- <b>CGAGTCT</b> - 625	
<b>Hb</b>	TGCCACCT-----AAAAGTATGCTATA <b>GTTGTGTCTTGTTCCTGCTGA</b> TCGAGTGT- 643	
<b>Hd</b>	----- <b>TATCTTGTTCCTGC</b> ----- <b>CGAGTCT</b> - 507	
Contig17	-----TTCTGC-----CGAGTCT- 625	
Contig11	-----TTCTTG-----CGAGTCT- 630	
Contig14r	-----TTCTTG-----CGAGTCT- 625	
Contig1	TGCAACCTG-----AAAGTATGCTACAGTT-GTGTCTTGTTCCTGG-----CGAGTCT- 674	
Contig15r	-----TATCTTGTTCCTGC-----CGAGTCT- 640	
Contig4a	TGCAACCTA-----AAATTATGCTAGTTGTTGCTTTGTTCTGT---CGAGTCTT 666	
Contig10	TGCAACCTCTGCAACCTAAAGTATGGTACGGTTGTCTTGTTCCTGC---CGAGTCT- 662	
Contiglar	TGCAACCT-----GAAAGTATGCTACAGTT-----CTTGCTGATCGAGTCT- 633	
Contig7	TGCAACCT-----GAAAGTATGCTACAGTT-----CTTGCTGATCGAGTCT- 634	
	* * ***** *	

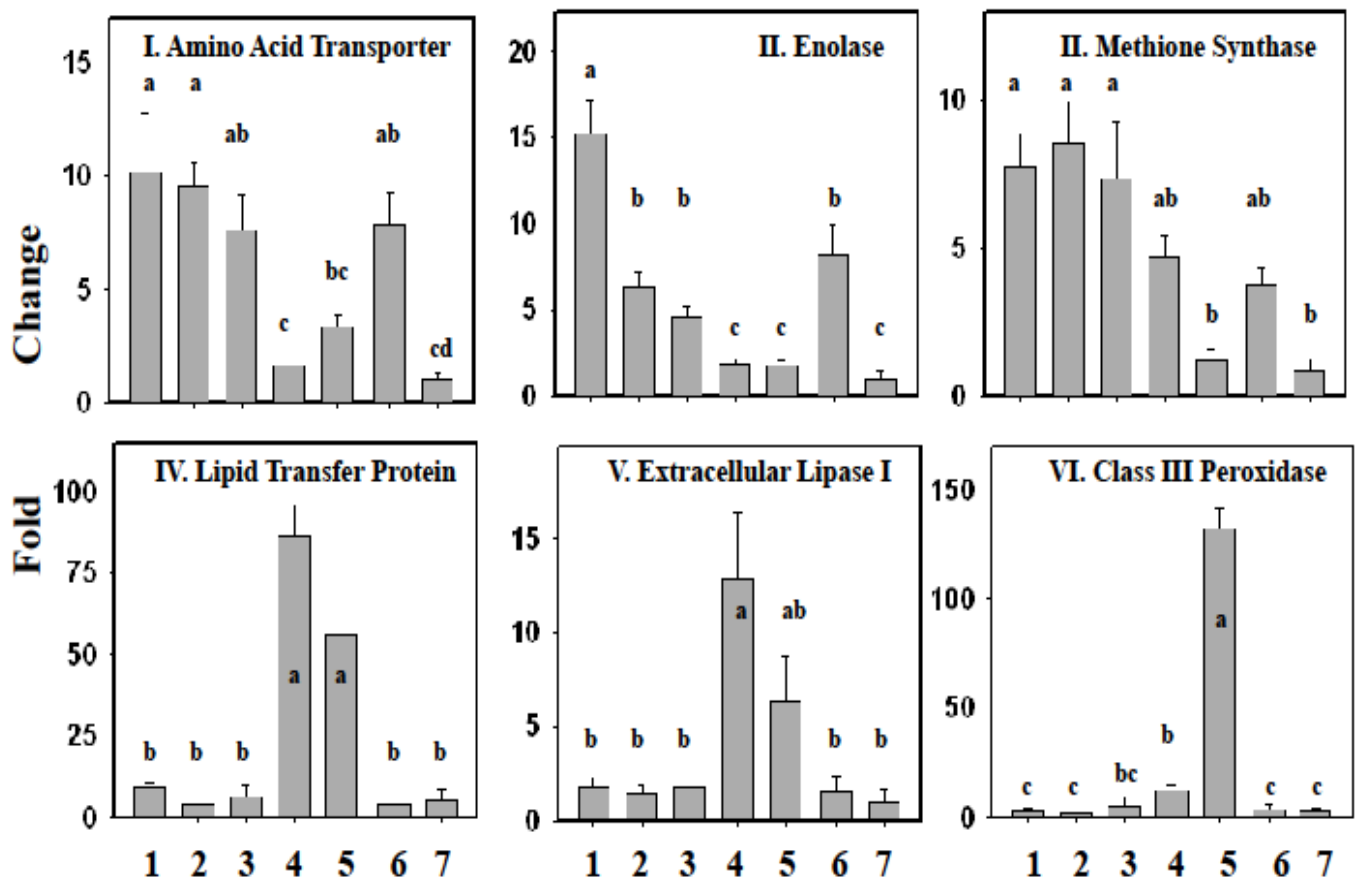
**Figure S9**



**Figure S10N**



**Figure S11**



**Figure S12**

